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(11) EP 0 921 395 A2

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication: 09.06.1999 Bulletin 1999/23

(51) Int Cl.6: G01N 33/48

(21) Application number: 98309266.9

(22) Date of filing: 12.11.1998

(84) Designated Contracting States:

AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU

MC NL PT SE

Designated Extension States:

AL LT LV MK RO SI

(30) Priority: 13.11.1997 US 65423 P

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(54) Assays for measurement of protein fragments in biological media

(57) This invention provides novel antibodies and engineered versions thereof and methodology for mon-

itoring biological media for protein fragments, especially collagen fragments resulting from collagenase deavage of type II collagen.

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Description

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FIELD OF THE INVENTION

[0001] This invention relates to methods for detecting protein fragments in biological media. More specifically, it relates to methods for quantitating collagen fragments resulting from collagenase cleavage of type II collagen.

BACKGROUND OF THE INVENTION

[0002] The physiological turnover of articular cartilage represents a fine balance between synthesis and degradation. It is a feature of normal growth and development and maintenance of cartilage in the adult. Net cartilage loss is a feature of rheumatoid arthritis and osteoarthritis. It is strongly associated with disability and a low quality of life. Cartilage destruction in rheumatoid arthritis and osteoarthritis is currently diagnosed based on combined clinical symptoms and radiological findings. Damage to articular cartilage occurs early in the disease, long before it can be detected radiologically; damage is detected radiologically only after there is extensive and probably irreversible cartilage loss. Therefore, it is of critical importance that clinicians have biochemical markers for early diagnosis of cartilage damage so therapy can be initiated early, before extensive damage is done.

[0003] Type II collagen constitutes the bulk of the fibrillar backbone of the cartilage matrix, just as type I collagen forms the fibrillar organization of the extracellular matrix of most other tissues such as skin, bone, ligaments and tendons. These collagens are composed of a tightly wound triple helix, which can only be cleaved by metalloproteinase collagenases to produce 3/4 and 1/4 length α -chain fragments that are identifiable by polyacrylamide gel electrophoresis.

[0004] The destruction of articular cartilage during arthritic disease is due, in part, to the degradation of the extracellular matrix, which is composed primarily of fibrillar type II collagen and aggregating proteoglycans. In articular cartilage, type II collagen fibrils are responsible for the tensile strength whereas the proteoglycans provide the compressive stiffness necessary for normal articulation and function. The precise mechanisms by which these connective tissue components are degraded are not fully understood. In mammals, an important mechanism involves the collagenases which are a group of enzymes capable of site-specific cleavage of helical (native) collagen.

SUMMARY OF THE INVENTION

[0005] This invention comprises a method for monitoring biological media for protein fragments, preferably, collagen fragments, said fragments resulting from collagenase cleavage of type II collagen which comprises; contacting said biological media with a capture antibody; said capture antibody being active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 1 and 2; and in a second step, contacting said biological media with a detection antibody; said detection antibody being active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 3 and 4; and finally, detecting the amount of collagen fragments bound to said capture and detection antibodies using standard techniques which are well known to those with ordinary skill in this art.

[0006] Those skilled in this art will recognize that the order of contacting the antibodies with the biological media may be reversed.

[0007] Therefore, in another aspect, this invention comprises a method for monitoring biological media for protein fragments which comprises;

detecting the amount of collagen fragments bound to said capture and detection antibodies; or contacting said biological media with a capture antibody; said capture antibody being active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 3 and 4; and contacting said biological media with a detection antibody; said detection antibody being active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 1 and 2; and detection antibodies.

[0008] In a preferred aspect, this invention provides a method for the detection of protein fragments which are collagen fragments generated by collagenase cleavage of articular cartilage and more particularly, a method wherein said protein fragments are generated from collagenase cleavage of type II collagen.

[0009] In yet another aspect, this invention provides a third method for monitoring biological media for collagen fragments generated from articular cartilage which comprises;

contacting said biological media with an antibody active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 3 and 4; and

detecting the amount of collagen fragments bound to said antibody.

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[0010] In a broader aspect this invention provides a method for monitoring biological media for protein fragments which comprises;

contacting said biological media with an antibody capable of recognizing and binding to protein fragments containing the sequences set forth in the Sequence Listing as SEQ ID NOS: 3 and 4; and detecting the amount of protein fragments bound to said antibody.

[0011] This invention provides a capture antibody which is a monoclonal antibody.

[0012] This invention also provides a detection antibody which is a monoclonal antibody.

[0013] This invention provides a monoclonal antibody, designated 9A4, which has the V_H sequence set forth in the Sequence Listing as SEQ ID NO: 5 and the V_L sequence set forth in the Sequence Listing as SEQ ID NO: 6.

[0014] This invention provides an antibody which is a genetically engineered antibody, and is related, but not necessarily identical in sequence to scFv 9A4 of p9A4ICAT7-1 (ATCC 98593), ATCC, American Type Culture Collection, 10801 University Blvd., Manassas, Virginia 20110-2209 USA, and p9A4IF-5 (ATCC 98592), which correspond to SEQ ID NOS: 7 and 8 set forth in the Sequence Listing, respectively.

[0015] This invention provides an antibody which is a genetically engineered antibody, and is related, but not necessarily identical in sequence to scFv 5109 of p5109CscFv7 (ATCC 98594), and as set forth in the Sequence Listing as SEQ ID NO: 9.

[0016] This invention provides a monoclonal antibody, designated 5109, which has the V_H sequence set forth in the Sequence Listing as SEQ ID NO: 10 and the V_L sequence set forth in the Sequence Listing as SEQ ID NO: 11.

[0017] This invention provides a method for detecting protein fragments employing an antibody, designated 5109, and which has the V_H sequence set forth in the Sequence Listing as SEQ ID NO: 10 and the V_L sequence set forth in the Sequence Listing as SEQ ID NO: 11.

[0018] In another aspect, this invention provides a hybridoma cell line that produces a monoclonal antibody that binds to peptides consisting essentially of the structure as set forth in the Sequence Listing as SEQ ID NO: 1 or SEQ ID NO: 2, the cell line having the identifying characteristics of ATCC HB-12436.

[0019] In another aspect, this invention provides a hybridoma cell line that produces a monoclonal antibody that binds to peptides consisting essentially of the structure as set forth in the Sequence Listing as SEQ ID NOS: 3 or 4, the cell line having the identifying characteristics of ATCC HB-12435.

[0020] In still another aspect, this invention provides an *E. coli* culture which produces a genetically engineered antibody related to 9A4 that binds to peptides consisting essentially of the structure set forth in the Sequence Listing as SEQ ID NOS: 1 or 2.

[0021] In another aspect, this invention provides an *E. coli* culture which produces a genetically engineered antibody related to 5109 that binds to peptides consisting essentially of the structure set forth in the Sequence Listing as SEQ ID NOS: 3 or 4.

[0022] In yet another aspect, this invention provides a bispecific antibody produced by hybridization of the antibodies 9A4 and 5109 wherein each half antibody recognizes its respective binding partner.

[0023] In another aspect, this invention provides a bispecific antibody which is a genetically engineered combination of antibodies 9A4 and 5109 produced by combining the V_L and V_H domains of the two antibodies in the form V_L(5109)-linker-V_H(5109)-linker-V_L(9A4)-linker-V_H(9A4) and equivalents thereof.

[0024] This invention further provides a method for monitoring collagen fragments in biological media which comprises: contacting said biological media with a bispecific antibody 9A4/5109 described above; and detecting the amount of collagen fragments bound to said antibody.

[0025] This invention also provides a bispecific antibody produced from genetically modified variants of antibodies 5109 and 9A4.

DETAILED DESCRIPTION OF THE INVENTION

[0026] Type II collagen is the structural protein that gives articular cartilage its tensile strength and shear resistance. It also provides the stuctural basis for the containment of proteoglycan that imparts compressive resistance and in doing so directly determines the form of the osmotically pressurized cartilage. Thus the structural integrity of type II collagen is a major determinant of the physical properties and the durability of articular cartilage.

[0027] The progressive failure of articular cartilage is one of the hallmarks of arthritic disease. Where that failure is based on changes in the type II collagen structure, it would be advantageous to have methodology to measure specificially the breakdown of type II collagen. Fragments of type II collagen from articular cartilage are released into the synovial fluid, lymph, blood and urine as type II collagen breaks down. Measurement of surviving fragments would

provide a method for monitoring type II collagen breakdown to detect the onset of arthritic disease and measure disease progression. Moreover, it would also be useful to measure the effect of therapy on type II collagen breakdown during disease.

[0028] A variety of methods have been utilized to monitor collagen breakdown. In mammalian tissues, collagenase appears to be the rate-limiting extracellular enzyme involved in breakdown of type II collagen (1). Collagenase fragmentation of collagen into a three quarters and one quarter piece was identified as early as 1967 (2,3) and there are currently three identified mammalian collagenases involved in breakdown of type II collagen (4,5). Other enzymes are involved in further fragmentation of type II collagen. Lysosomal breakdown of the fibrillar collagens is known in bone and liver (6,7). Since collagen is one of the few proteins characterized by a high hydroxyproline content, measurement of urinary hydroxyproline has been examined as a measure of collagen tumover (8). However, since type I and type III collagens are found in large amounts in skin, bone, and connective tissues in general, the method has not been found useful for measuring breakdown of either a particular type of collagen or of collagen from a particular body compartment, e.g., bone, and is of no value for monitoring type II collagen since it provides only an extremely small portion of the daily urinary secretion of hydroxyproline (8). Therefore efforts to monitor breakdown of collagen or collagen itself have focused on immunological methods.

[0029] Antibodies can discriminate collagen fragments unique to a collagen type and cleavage site and potentially monitor the specific mode of collagen breakdown (9). Polyclonal and monoclonal antibodies have been prepared against type I and III collagen or their fragments; and assays have been prepared for the collagen breakdown products of type I and type III collagens, for example, see Eyre (10). Relatively few methods have been reported to develop antibodies against breakdown products of type II collagen.

[0030] Polyclonal and monoclonal antibodies have been prepared against type II collagen (9,11-13). These antibodies have been utilized for the detection of intact type II collagen rather than the quantitative determination of collagen fragments. Eyre, however, (14) has prepared monoclonal antibodies against type II collagen fragments containing the crosslinking residues. He developed an assay for breakdown of type II collagen based on the crosslink fragment containing a type II collagen specific sequence similar to his issued patent (10). Dodge and Poole prepared polyclonal antibodies against denatured type II collagen that were unreactive with other collagens (15,16). The epitope was sequenced and later Hollander and Poole (17,18,19,) prepared a competitive antibody assay against the type II collagen fragments having the sequences set forth in the Sequence Listing as SEQ ID NOS: 12 and 13 using a monoclonal antibody. Hollander and Croucher (20) also made a capture Elisa using antibodies directed against peptides outlined in the Sequence Listing as SEQ ID NO: 67, 68 or 69. Billinghurst et al. (21) have prepared polyclonal antibody against the collagen cleavage site necepitope of type II collagen (having the sequence set forth in the Sequence Listing as SEQ ID NO: 2) and have prepared a competitive type II collagen assay. Srinivas, Barrach, and Chichester (22-24) have prepared multiple monoclonal antibodies for a type II collagen assay using the cyanogen bromide fragments of type II collagen as antigen. Although the epitopes reactive with the antibody have not been identified (25), they too are able to assay type II collagen.

Capture Elisa's are a reliable method for obtaining high specificity since it is based on two antibodies coordinately recognizing two different amino acid clusters in the same molecule. As the antibody binding site contains as few as six amino acids, sequences as small as 15 to 20 amino acids can be recognized by two antibodies in a capture assay. Competitive assays use a single antibody and thus allow measurement of polypeptides as small as 6-8 amino acids, but they lack the specificity of the dual antibody measurement. In addition, competitive assays lack the sensitivity of capture Elisa's which often have a 100-10,000 times lower limit of detection. Thus capture Elisa's provide a preferred method for measuring metabolic fragments of proteins. For example, a capture Elisa has been used to measure breakdown fragments of the structural protein elastin in blood (26). A capture Elisa has been used to measure the 21 amino acid biologically active peptide endothelin (27). A pair of capture Elisa's have been used to measure different metabolic fragments of the 28 amino acid glucagon peptide (28). Based on such results, a minimal 22 amino acid peptide fragment was selected that could be used to construct a capture Elisa to measure collagenase dependent metabolism of type II collagen.

[0031] This invention provides two types of assays of type II collagen metabolism. Both assays are based on an antibody (polyclonal, monoclonal or genetically engineered antibody) against a defined sequence of type II collagen against which antibodies have not been previously prepared. The sequence is rich in acidic residues, i.e., the sequence set forth in the Sequence Listing as SEQ ID NO: 3 from which a deletion has been made at the C-terminus by 1 residue. As no mammalian extracellular aspartyl or glutamyl endopeptidases have been described, collagen fragments rich in acidic residues should survive further metabolism and be available for measurement in body fluids. Antibody against those residues or a collagen fragment containing those residues would provide a general method of detection of type II fragments in body fluids independent of the method of generation of the collagen metabolite. This invention provides monoclonal antibody 5109 and genetically engineered variants of 5109 that are specific for the type II collagen sequence and bind to collagen fragments containing the sequence.

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[0032] The first assay is a general method for assessing the breakdown of type II collagen. This assay provides a

general competitive method for quantitating the amount of the sequence set forth in the Sequence Listing as SEQ ID NO: 3 from which a deletion has been made at the C-terminus by 1 residue, and its closely related congeners.

[0033] For the second assay, an additional antibody was made against the sequence set forth in the Sequence Listing as SEQID NO: 14. There is a free C-terminal carboxyl group on the glycine (residue 9 of SEQID NO: 14). This sequence is obtained when collagenase cleaves type II collagen and therefore it is classified as a necepitope, i.e., it is not present in the native sequence (the sequence set forth in the Sequence Listing as SEQ ID NO: 15, continues -GPOGPQG/ LAG-where collagenase cleaves at the vertical bar), but arises when collagen is cleaved by collagenase. Polyclonal antibodies against that sequence have been previously reported (22). Monoclonal antibody 9A4 and genetically engineered derivatives therefrom react with the necepitope sequence set forth in the Sequence Listing as SEQ ID NO: 2. but fail to react with uncleaved type II collagen or type I collagen. While the necepitope sequence is unique to type II collagen when cleaved by collagenase, a homologous and weakly cross-reactive sequence is generated in type I collagen when cleaved by collagenase (SEQ ID NO: 16 as set forth in the Sequence Listing). This is also true for type III collagen; it generates the weakly cross-reactive sequence set forth in the Sequence Listing as SEQ ID NO: 2, when cleaved by collagenase. Thus the necepitope antibody lacks full specificity for type II collagen and would fail to selectively detect cleavage of type II collagen by collagenase if used alone. However, when antibody 5109 and antibody 9A4 are combined in a sandwich assay the two antibodies together can selectively detect type II collagen metabolites generated by collagenase. Moreover, an advantage that the sandwich type of assay provides in this invention is a 100 fold lower limit of detection compared to a simple competitive assay based on 9A4 alone. The sandwich assay format with the antibodies described in this invention thus provides a unique method for monitoring type II collagen metabolism by collagenase in normal and pathological conditions, which has not been previously described.

[0034] Alone, antibody 9A4 is a novel monoclonal antibody that has use for the detection of collagenase cleaved fragments of type I or type II or type III collagen, so long as it is not necessary to distinguish the collagen type.

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DEFINITIONS

- [0036] Immunoglobulin (Ig): A natural tetrameric protein composed of two light chains of circa 23 kD and two heavy chains of circa 53-70 kD depending on the amino acid sequence and degree of glycosylation. Multimers of the tetrameric protein are also formed (IgM and IgA). There are two classes of light chains, kappa (κ) and lambda (λ), and several classes of heavy chains gamma (γ), mu (μ), alpha (α), delta (δ) and epsilon (ε). There are also subclasses. Each chain, whether a light chain or heavy chain, is made up of two parts. The first part, beginning from the N-terminus of either chain, is called the variable domain. The C-terminal half of the light chain is called the constant region of the light chain and it is the primary determinant whether the light chain is a κ or λ type. The constant region of the heavy chain comprise circa the C-terminal three-fourths of the heavy chain and determines the class of the immunoglobulin molecule (IgG₁, IgM, etc), i.e. a X heavy chain corresponds to an IgG and a μ heavy chain corresponds to an IgM, ect.
- [0037] V_L and V_H: The amino acid sequence of the variable domain of the light chain (V_L) and the variable domain of the heavy chain (V_H) together determine the binding specificity and the binding (kD) constant of the immunoglobulin molecule. The variable domain comprises circa half the length of the light chain and circa a quarter of the length of the heavy chain and for both chains, begins at the N-terminus of the chain. The variable regions each contain three (3) hypervariable segments known as the complementarity determining regions or CDRs.
- [0038] CDR and FR: Each variable domain, V_L or V_H, is comprised of three CDRs: CDR1, CDR2 and CDR3. The intervening sequence segments before, between and after the CDRs are known as framework segments (FR). Each V_L and V_H is comprised of four FR segments FR1, FR2, FR3 and FR4.
 - [0039] V_{κ} and V_{λ} : The V_L domain is either κ or λ , depending on which constant region (C_{κ} or C_{λ}) is used during the productive rearrangement of the light chain (VJC $_{\kappa}$ or VJC $_{\lambda}$).
- [0040] Antibody: Antibodies are specific immunoglobulin molecules produced by B-cells of the immune system in response to challenges by proteins, glycoproteins, virus cells, chemicals coupled to carriers, and other substances. An antibody is simply an immunoglobulin molecule for which its binding partner is known. The substance to which the antibody binds is called an antigen. The binding of such antibodies to its antigen is highly refined and the multitude of specificities capable of being generated by changes in amino acid sequence in the variable domains of the heavy and light chains is remarkable.
- 55 [0041] Polyclonal antibody: Normal immunization leads to a wide variety of antibodies against the same antigen. Although each B lymphocyte normally produces one immunoglobulin molecule of a defined amino acid sequence, in an immune response, many B lymphocytes are stimulated to make immunoglobulin molecules that react with the antigen, i.e., antibodies. These different antibodies are characterized by different amino acid sequences in the variable

regions of the immunoglobulin molecule which result in differences in the fine specificity and affinity of binding. Such antibodies are called polyclonal antibodies to emphasize the variety of binding specificities and binding constants which arise from the variety of amino acid sequences found in the different immunoglobulin molecules utilized in the immune response.

[0042] Monoclonal antibody (MAb): A B lymphocyte producing a single antibody molecule can be hybridized with an immortal B lymphocyte cell line, i.e., a myeloma, to derive an antibody producing immortal cell line, i.e., a hybridoma. The hybrids thus formed are segregated into single genetic strains by selection, dilution, subcloning, and regrowth, and each strain thus represents a single genetic line. It produces a single antibody of a unique sequence. The antibody produced by such a cell line is called "monoclonal antibody" or MAb, referencing its pure genetic parentage and differentiating it from polyclonal antibody, produced from a mixed genetic background, i.e., multiple B cells. Because a MAb is a pure chemical reagent it gives consistent, uniform results in immune tests. Moreover, because the MAb is produced by an immortal cell line, reagent supply is not limiting. For these reasons, a MAb is much preferred over polyclonal antibodies for diagnostic purposes.

[0043] Genetically engineered antibody: As the binding specificity of an antibody resides in the variable regions of the light and heavy chains, antibodies can be genetically engineered to change or remove the constant regions and, if done properly, it can result in an antibody molecule with different properties and molecular weight, but with the same or very similar antigen binding properties. For example, the V_L and V_H genes can be cloned and assembled (or V_H and V_L) with an appropriate linker between them. Such a new genetically engineered molecule is called a single chain antibody (abbreviated scFv) and typically has a molecular weight of 25-28 kD depending on the design of the linker and the addition of other sequences to help in purification, stability, trafficking, detection, etc. Multimers of the single chain antibody can also be made by appropriate use of linkers in which the order of each V_H, V_L pair may vary. In addition, some changes in the amino acid sequence of the V_L and V_H region can be made that retain desirable antigen binding properties. It can be seen that an infinite variety of genetically engineered antibodies can be derived from the original antibody sequence which retain binding specificity to the antigen, but which are tailored to fulfill specific requirements. Other examples of genetically engineered antibodies include, but are not limited to: Fab, F(ab')₂, chimeric antibodies, humanized antibodies, etc. For a review, see Winter G and Milstein C, "Man-made Antibodies", Nature 1991; 349, 243-299.

[0044] <u>Bispecific antibody:</u> Normally an IgG antibody has two identical light chains and heavy chains. There are therefore two identical antibody binding sites in the immunoglobulin molecule. By contrast, a bispecific antibody is a single immunoglobulin molecule which has two specificities. It can be made by fusion of two monoclonal antibody producing hybridoma cell lines, where each hybridoma has a different antigen specificity, and selection for a cell line (a quadroma) that produces an antibody whose composition is a tetramer composed of one light chain and one heavy chain from each hybridoma fusion partner. The antibody produced by the quadroma has only one light and one heavy chain of each parental specificity and has one binding site for each heavy/light chain pair and is bispecific, i.e., it has two binding sites of different specificities. A bispecific antibody can also be made by genetic engineering. It can comprise the V_L linker V_H of one antibody linked through an additional linker to a V_L linker V_H of another antibody molecule. The order of V_L and V_H can be altered, but the end result is a bispecific antibody.

[0045] Epitope: Depending on the size, structure and conformation of the antigen, an antibody may bind only to a small part of the entire structure. The part of the antigen molecule to which the antibody binds is called its epitope. Different antibodies may be mapped to different epitopes on the same antigen.

[0046] Neoepitope: The antigen may have an epitope which is hidden so that it cannot bind to a specific antibody. However, a conformational change in the antigen may cause the appearance of the epitope by unfolding or uncovering part of the surface of the molecule. This now allows the antibody to bind to the epitope. In another aspect, the action of an enzyme on the antigen may cause the appearance of a new epitope to which the antibody can bind. For example, after cleavage by a proteolytic enzyme, new N-terminal and new C-terminal sequences are generated. Because the epitope is not observed in the parent molecule and because, after some change in the parent molecule, the epitope is revealed and now can bind antibody, it is called a necepitope.

[0047] <u>Biological media:</u> This may be defined as any biological fluid that might contain the antigen and be of interest to assay by this procedure. These include: blood, synovial fluid, urine, spinal fluid, bronchiolar lavage fluid, lymph, the vitreous humor of the eye, extracts of tissues, tissue culture supernatants, extracts of cartilage, etc., Biological media need not be limited to human samples, but may also be obtained from a similar variety of animal media (mouse, rat, hamster, guinea pig, dog and bovine have been tested) in a fashion similar to the examples above.

[0048] Immunoassay: An assay for a substance (complex biological such as a protein or a simple chemical) based on using the binding properties of antibody to recognize the substance which may be a specific molecule or set of homologous molecules. The assay may involve one or more antibodies.

[0049] <u>Direct assay:</u> The antibody binds directly to an antigen such as in a biological specimen (cells, tissues, histological section, etc.) or to antigen adsorbed or chemically coupled to a solid surface. The antibody itself is usually labeled to enable the determination of the amount of antibody bound to the antigen. Alternatively, the antibody (now

termed primary antibody) is detected with a secondary labeled antibody that will demonstrate that binding of the primary antibody had occurred.

[0050] Competitive assay: An assay based on the binding properties of a single antibody molecule. Typically, a labeled antigen is used to compete with an unknown antigen and the amount of unknown antigen is determined in terms of how much of the labeled antigen is displaced by the unknown antigen. The label may be radioactive, optical, enzymatic, florescent polarizing, florescent quenching, or other label. The antibody may be monospecific or bispecific. [0051] Sandwich assay: This is a double antibody assay in which both antibodies bind to the antigen, forming a trimeric immune complex or sandwich containing the two antibodies with the antigen between them. One antibody is utilized to localize the immune complex to the detection surface or chamber. This antibody is termed the capture antibody. The other antibody bears a label that will allow the immune complex to be detected. It is called the detection antibody. If an immune complex is not formed (no antigen is present), then the capture antibody is unable to bring the detection antibody to the detector. If antigen is present, then an immune complex will form and the capture antibody will be joined with the detection antibody such that the amount of detection antibody in the immune complex is quantitatively related to the amount of antigen present.

[0052] The assay can be formatted in many ways. For example, the capture antibody can be chemically coupled to a solid surface, or non-specifically adsorbed to a surface, attached via biotinylation to an avidin-like molecule, eg. avidin, streptavidin, neutravidin, etc., streptavidin or avidin-coated surface, coupled to magnetic particles or beads as a means of localizing the immune complex to the measurement device.

[0053] The detection antibody may be radiolabeled, or it may have a variety of possible enzymatic amplification systems such as horse radish peroxidase (HRP), alkaline phosphatase (AP), urease, etc., when formatted as an Elisa (Enzyme-linked immune assay). It may have an electrochemical, an optical, a fluorescent or other detection method to determine the amount of detection antibody in the immune complexes.

[0054] It may immediately be seen that many examples can be derived in which the two antibodies are paired in a sandwich assay using a variety of methods to capture the immune complex in a detection device and a variety of detection systems to measure the amount of immune complex.

[0055] Molecular biology techniques: Because the nucleotide sequences of V_H and V_L -encoding regions are now provided for the antibodies of the present invention, a skilled artisan could *in vitro* produce a complete gene coding for the V_H and V_L regions and a completely functional antibody. It can be produced as an immunoglobulin molecule of any given class with constant regions of the heavy chain and light chain added or it can be produced as a scFv with V_H and V_L joined by a linker with tags added as appropriate. The constructed gene may be engineered by conventional recombinant techniques, for example, to provide a gene insert in a plasmid capable of expression. Thereafter, the plasmids may be expressed in host cells where the host cells may be bacteria such as *E. coli* or a *Bacillus* species, yeast cells such as *Pichia pastoris* or in mammalian cell lines such as Sp2/0, Ag8 or CHO cells.

<u>Abbreviations</u>

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[0056] Nucleic acids, amino acids, peptides, protective groups, active groups and similar moieties, when abbreviated are abbreviated according to the IUPACIUB (Commission on Biological Nomenclature) or the practice in the fields concerned. The following are examples.

Standard Ab	Standard Abbreviations				
HPLC SDS-PAGE PCR Oligo RT	High pressure liquid chromatography Sodium dodecylsulfate polyacrylamide gel electrophoresis Polymerase chain reaction Oligonucleotide Room temperature, circa 22°C.				
Reagents:					
EDTA SDS TW-20 NFDM DPBS Bt HAT HT	S Sodium dodecylsulfate 7-20 Tween-20 DM Non-fat dry milk BS Dulbecco's phosphate buffered saline Biotinylated T Hypoxanthine, aminopterin, thymidine containing media				

(continued)

Standard Ab	Standard Abbreviations				
Reagents:	Reagents:				
HRP	HRP Horseradish peroxidase				
Immunoglobulin-like molecules or chains					
V _H or VH V _L or VL scFv	Variable region of the heavy chain Variable region of the light chain Single chain antibody containing a V _L and V _H				
Nucleic Acid	Nucleic Acids				
RNA DNA cDNA mRNA	Ribonucleic acid Deoxyribonucleic acid Complimentary DNA Messenger RNA				

Nucleic acid bases
Purines Pyrimidines
A: Adenine
G: Guanine
C: Cytosine
U: Uracil

Amino Adds-Single letter codes: Three letter codes: Full names. G Gly glycine Val valine Leu leucine Α Ala alanine lle isoleucine s Ser serine D Asp aspartic acid Κ Lys lysine R arginine Arg н His histidine F Phe phenylalanine Υ tyrosine Tyr T Thr threonine С Cys cysteine М Met methionine Ε Glu glutamic acid W Trp tryptophan Ρ Pro proline 0 hydroxyproline Нур Ν asparagine Q Asn Gln glutamine

Abbreviations used in the Sequence Listing

[0057] The sequence listing has been prepared in accordance with WIPO Standard ST.25. [0058] The most important abbreviations are shown in the following table:

<110>	Applicant name			
<120>	Title of invention			
<160>	Number of SEQ ID NOs			
<210>	SEQ ID NO: x			
<211>	Length			
<212>	Туре			
<213>	Organism			
<400>	Sequence			

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Example 1

Generation and characterization of monoclonal antibody 9A4,

[0059] Balb/c mice (Jackson Laboratories, Bar Harbor, ME) were immunized initially with the peptide having the sequence set forth in the Sequence Listing as SEQ ID NO: 17 (Anaspec, San Jose, CA) covalently linked to the KLH maleimide (Pierce Chemical, Rockford, IL) and administered in complete Freund's adjuvant (DIFCO Detroit, MI). The mice were boosted monthly for about 5 months using incomplete Freund's adjuvant (DIFCO, Detroit, MI) until the titers were 1:100,000. Mice were boosted i.v. 10 days prior to fusion. Splenocytes were collected and fused with a non-lg secreting cell-line derived from P3X63Ag8.653 (American Type Culture Collection ATCC, Bethesda, MD) using 50% PEG-1450 (ATCC). They were plated at 10⁶ cells/well in 96 well microtiter plates in HAT media (Sigma, St. Louis, MO) with 15% fetal calf serum (Hyclone, Provo, Utah). Ten days later the wells were screened by a primary Elisa. For identification of positive antibody producing wells, 10 nglmL biotinylated peptide (Bt-AEGPPGPQG) [biotinylated on residue of 1 of SEQ ID NO: 14] was added to streptavidin (10 μg/mL) coated plates (Pierce Chemical) and 2 μL of each hybridoma supernatant added to 100 μL of DPBS (Gibco, Grand Island, NY) with 0.05% TW-20 (Sigma). Elisa positive wells were detected by rabbit anti-mouse IgG-HRP (Jackson Immuno Research, West Grove, PA). [0060] Positive wells in the Elisa were subjected to a second round of selection on the BIAcore. Wells were sought which produced antibodies having slow off rates on the BIAcore as determined using BIAevaluation version 2.1 software (Pharmacia Biosensor, Piscataway, NJ). Streptavidin (Pierce Chemical) at 100 µg/mL was conjugated using the Pharmada Amine Coupling Kit (Pharmacia Biosensor) to carboxylated dextran-coated biosensor chips (Pharmacia Biosensor) at pH 4.0 using a flow rate of 5 µL/minute for 35 minutes. Typically, 2000 RU was added. Peptide (100 ng/mL)

sor) at pH 4.0 using a flow rate of 5 μL/minute for 35 minutes. Typically, 2000 HU was added. Peptide (100 ng/mL) biotinylated on residue of 1 of SEQ ID NO: 14 as set forth in the Sequence Listing, was passed over the streptavidin chip at a flow rate of 100 μl/minute for 10 seconds. Candidate supernatants containing antibody were passed over the chip (2 μL/minute for 30 seconds) and the amount of added antibody noted. The buffer was changed to HBS (Pharmacia Biosensor), and the dissociation rate noted for the next 80 seconds. The chip was cleaned with 0.1 N HCI for 30 seconds between each run to remove residual antibody and to clear any nonspecific binding. The off-rates were determined using BlAevaluation kinetic analysis software version 2.1. Clones with the slowest off rates were selected for further analysis. These clones included 9A4, 11F2 and 3H10.

[0061] Further characterization of these clones was performed as follows: Four preparations consisting of type I collagen, type I collagen cleaved by collagenase, type II collagen, and type II collagen cleaved by collagenase were each coupled to a separate flow cell on a BIA 2000 instrument. They were conjugated using the Pharmada Amine Coupling Kit (Pharmacia Biosensor) to carboxylated dextran coated biosensor chips (Pharmacia Biosensor) at pH 4.0 using a flow rate of 5 μ L/minute for 35 minutes. The four flow cells added 8000, 7000, 4000 and 4000 RU, respectively. The cells were washed with 0.1 N HCI to clean them of any uncoupled material and to clean them of any residual antibody between runs. All antibody preparations were purified by Protein G chromatography (Pharmacia Biotechnology, Piscataway, NJ) and were run at 10 μ g/mL. The total binding to each of the four surfaces was recorded. Antibody 9A4 was selected because it showed selective binding to collagenase-cleaved type II collagen (type I = 11 RU; type II = 280 RU) and lacked significant binding to uncleaved collagen (type I = 3 RU; type II = 6 RU).

[0062] After three rounds of subcloning by limiting dilution in HT media (Sigma) with 5% fetal calf serum (Hyclone), a stable 9A4 monoclonal hybridoma was obtained. It has been deposited with the American Type Culture Collection as ATCC - HB-12436.

Example 2

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Generation and characterization of monoclonal antibody 5109.

[0063] Balb/c mice were immunized initially with the peptide having the sequence set forth in the Sequence Listing as SEQ ID NO: 18 (Anaspec, San Jose, CA) covalently linked to the KLH maleimide (Pierce Chemical) and administered in complete Freund's adjuvant (DIFCO). The mice were boosted monthly for about 5 months using incomplete Freund's adjuvant (DIFCO) until the titers were 1:100,000. Mice were boosted i.v. 10 days prior to fusion. Splenocytes were collected and fused with a non-lg secreting cell-line derived from P3X63Ag8.653 cells (ATCC) using 50% PEG-1450 (ATCC). They were plated at 10⁶ cells/well in 96 well microtiter plates in HAT media (Sigma) with 15% fetal calf serum (Hyclone). Ten days later the wells were screened by Elisa. For identification of positive antibody producing wells, 10 ng/mL biotinylated peptide (biotinylated on residue 1 of SEQ ID NO: 19 as set forth in the Sequence Listing) was added to streptavidin (10 μg/mL) coated plates (Pierce Chemical) and 2 μL of each hybridoma supernatant added to 100 μL of DPBS with 0.05% TW-20 (Sigma). Elisa positive wells were detected by rabbit anti-mouse IgG-HRP (Jackson ImmunoResearch).

[0064] Positive wells were subjected to a second round of selection on the BIAcore for those which had antibodies

giving the slowest off-rates on the BIAcore. Streptavidin (Pierce Chemical) at 100 μg/mL was conjugated using the Pharmacia Amine Coupling Kit (Pharmacia Biosensor) to carboxylated dextran coated biosensor chips (Pharmacia Biosensor) at pH 4.0 using a flow rate of 5 μl/minute for 35 minutes. Typically, 2000 RU was added. Peptide (100 ng/mL), biotinylated on residue 1 of SEQ ID NO: 19 as set forth in the Sequence Listing, was passed over the streptavidin chip at a flow rate of 100 μL/minute for 10 seconds. Supematants containing antibody were passed over the chip (2 μL/minute for 30 seconds) and the amount of added antibody noted. The buffer was changed to HBS (Pharmacia Biosensor), and the dissociation rate noted for the next 80 seconds. The chip was cleaned with 0.1 N HCl for 30 seconds between each run to remove antibody and to get rid of any nonspecific binding. The off-rates were determined using the BIAevaluation software version 2.1. Clones with slow off-rates were sought. MAb 5109 was selected for further analysis.

[0065] Four preparations consisting of type I collagen, type I collagen cleaved by collagenase, type II collagen, and type II collagen cleaved by collagenase were each coupled to a single channel of a four channel BIAcore. These were conjugated using the Pharmada Amine Coupling Kit (Pharmacia Biosensor) to carboxylated dextran coated biosensor chips (Pharmacia Biosensor) at pH 4.0 using a flow rate of 5 µL/minute for 35 minutes. The four flow cells added 8000, 7000, 4000 and 4000 RU, respectively. The cells were washed with 0.1 N HCI to clear them of any uncoupled material and to clear them of any specific materials between runs. All antibody preparations were purified by Protein G chromatography (Pharmacia Biotech), and run at 10 µg/mL. The total binding to each of the four surfaces was recorded. Antibody 5109 was selected because it showed selective binding to collagenase cleaved collagen (cleaved type I = 23 RU; cleaved type II = 173 RU), but lacked significant binding to uncleaved collagen (type I = 23 RU; type II = 15 RU). After nine rounds of subcloning by limiting dilution in HT media (Sigma) with 5% fetal calf serum (Hyclone), a stable 5109 monoclonal hybridoma was obtained. It has been deposited with the American type culture collection as ATCC - HB-12435.

Example 3

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Description of a sandwich assay using 9A4 as the capture antibody and monoclonal 5109 as the detection antibody.

[0066] Monoclonal antibody 9A4 (the capture antibody) was added to Nunc Maxisorp (VWR, Boston, MA) 96-well plates with 9A4 at 10 µg/mL in 0.05M sodium borate buffer, pH 8.5 using 100 µL/well (except for control wells numbered 4, 5 and 6, see Table 1) and incubated for 18-48 hours at 4° C.

[0067] The plate was washed three times with DPBS with 0.05% TW-20 (Sigma), (DPBS/TW-20); 200 μL/well was used.

[0068] Wells in the plate were blocked with 1% non-fat dry milk (NFDM) dissolved in DPBS (NFDM DPBS) prepared freshly, i.e., held on ice for no more than the day of use, using 100 µL/well incubated for 1 hour at RT.

[0069] The blocking solution was discarded, the wells rinsed one time with 200 µL of DPBS/TW-20.

[0070] Peptide 130 was diluted in 0.1% NFDM DPBS to concentrations shown in Table 1. Peptide 130 has the sequence set forth in the Sequence Listing as SEQ ID NO: 20 and was synthesized and purified by Anaspec Inc (San Jose, CA).

[0071] The dilutions of peptide 130 (SEQ ID NO: 20), the specimens at appropriate dilutions, and the controls were placed into the specified wells of the microtiter plate as shown in Table 1.

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Table 1. Outline of the microtiter plate and antibody coating scheme

pep130 (SEQ ID NO: 20) ng/mL

рер	2	1.33	0.889	0.59	0.4	0.26	0.18	0.12	0.08	0.05	0.03	0.02
130	*	89	**	*	-	R		11	**	Ħ	*	н
	smpl	smpl	smpl	smpl	smpl	smpl	smpl	smpl	smpi	smpl	smpl	smpl
	₩	**	*		"	**	**		М	H		-
	**	*	**	*	-	*	н		H		"	-
	**	*	*	"	-	н	in .	*	W	н	-	-
	17	**	*	"	-	11	н	*	**	×	-	
cntrls.	1	1	2	2	3	3	4	4	5	5	6	6

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Table 2.

	Additions to the control wells						
	Controls:		Biotinylated	Anti-biotin antibody HRP-labelled			
	9 A 4	130	5109				
1	+	-	-	+			
2	+	+	•	+			
3	+	-	+	+			
4	-	+	+	+			
5	-	+	•	+			
6	-	<u> </u>	+	+			

[0072] The wells were washed three times with 200 µL/well of DPBS TW-20.

[0073] Biotin-conjugated MAb 5109 (Bt-5109) was added to all peptide 130 (SEQ ID NO: 20) containing wells, all sample wells, and all control wells except 1,2, and 5. 5109-Bt (100 µL/well) at 1 µg/mL in 0.1% NFDM DPBS was added to each well and the plate incubated for 40 min at 37° C.

[0074] Note: MAb 5109 was biotinylated using 37 μg of biotin-N-hydroxysuccinamide (Pierce Chemical) per mg of MAb 5109 for 2 hrs and then dialyzed overnight using a 10 kD cutoff dialysis cassette (Pierce Chemical).

[0075] The wells were washed three times with 200 μ L/well of DPBS TW-20.

[0076] Mouse monoclonal anti-biotin antibody conjugated with HRP (Jackson ImmunoResearch) was diluted 1/5000 in 0.1% NFDM DPBS and 100 μ L/well was added to all wells and incubated for 30 minutes at RT.

[0077] The wells were washed three times with 200 µL/well with DPBS TW-20.

[0078] 100 µL/well of 1-step Turbo (ready to use 3,3',5,5'-tetramethyl benzidine; Pierce Chemical) was added to each well and incubated at RT for approximately 10 minutes. Color development was stopped with 2N H₂SO₄. The results were read on a spectrophotometer at 450 nm.

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Table 3. Standard curve data for 9A4 capture/Bt-5109 detection sandwich assay

E311	pep130	(SEQ ID NO: 20)			lin	lin regre	ssion
	ng/mL	nM	nM	OD450	regr	slope	intercept
1	10	5.88	0.769	0.78		0.029	0.024
2	5	2.94	0.468	0.74			
3	2.5	1.47	0.167	0.698			
4	1.25	0.735	-0.134	0.623	0.618		
5	0.625	0.368	-0.435	0.483	0.474		
6	0.313	0.184	-0.736	0.326	0.331		
7	0.156	0.092	-1.037	0.151	0.187		

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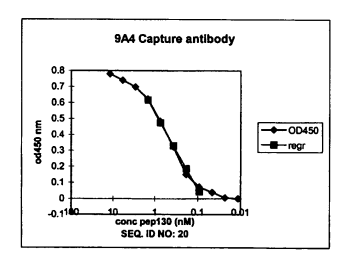
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0.078 0.046 0.072 8 -1.338 0.044 9 0.039 0.023 -1.639 0.039 10 0.020 0.011 -1.94 0.005 0.010 11 0.006 -2.241 -5E-04 12 0 0 -0.006

[0079] From the concentrations of peptide 130 and the resulting optical density reading at 450 nm (Table 3), a standard curve was constructed (Figure 1). Other appropriate peptides or collagen fragments can be substituted to prepare a standard curve analogous to Figure 1. The units were expressed in terms of molar equivalents of standard. In this case, the units were nM equivalents of peptide 130 (SEQ ID NO: 20).

[0080] Over the linear portion of the curve, a regression line was used to fit the data. In the given case, the standard curve was linear between 0.735 nM and 0.46 nM when the concentrations are given in the log scale (as in the example, Figure 1) and the regression curve is obtained using just the linear portion of the curve. For samples that fall outside of the linear portion, the concentrations can be read off the graph or the samples may be diluted to fall within the standard portion of the curve, or they may be below the limit of detection.

Figure 1. Standard curve of peptide 130 (SEQ ID NO: 20) and resulting optical density reading determined 9A4 Capture/Bt-5109 Sandwich Elisa.



[0081] The regression between log (nM) and OD450 gives a slope of 0.029 OD450/log(nM) and an intercept of 0.024 OD450. When unknown samples are run, the calibration curve can be used to determine of concentration of colla-

genase-generated type II collagen fragments from the optical density of the sample. The following equation can be used:

Log(Concentration) = (Sample OD450 - Intercept)/Slope

Sample:

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[0082] In the given case, an unknown sample of synovial fluid had an OD450 of 0.229. Thus

Log(Concentration) = (Sample OD450 - 0.024)/0.029 = -0.949

Taking the anti-log, the concentration of fragment in synovial fluid = 0.112 nM

15 Example 4

Description of a sandwich assay using monoclonal antibody 5109 as the capture antibody and 9A4 as the detection antibody.

20 [0083] Monoclonal antibody 5109 (the capture antibody) was added to Nunc Maxisorp (VWR, Boston, MA) 96-well plates with 5109 at 10 μg/mL in 0.05 M sodium borate buffer, pH 8.5, using 100 μL/well (except for control wells numbered 4, 5 and 6, see Table 4) and incubated for 18-48 hours at 4° C.

[0084] The plate was washed three times with 200 μ L/well of DPBS TW-20.

[0085] Wells in the plate were blocked with 1% (NFDM) dissolved in DPBS prepared freshly using 100 μL/well incubated for 1 hour at room temperature.

[0086] The blocking solution was discarded, and the wells rinsed one time with 200 μL of DPBS/TW-20.

[0087] Peptide 130 (SEQ ID NO: 20) was diluted in 0.1% NFDM DPBS to concentrations shown in Table 4. Peptide 130 has the sequence set forth in the Sequence Listing as SEQ ID NO: 20 and was synthesized and purified by Anaspec Inc (San Jose, CA).

[0088] The dilutions of peptide 130 (SEQ ID NO: 20), the unknown specimens at appropriate dilutions, and the controls were placed into the specified wells of the microtiter plate as shown in Table 4.

Table 4. Outline of the microtiter plate and antibody coating scheme

pep130 (SEQ ID NO: 20) (ng/mL) 1.33 0.889 0.59 0.4 0.26 pep 0.18 0.12 0.08 0.05 0.03 0.02 130 smpl . M " * ** * H Ħ ** cntrls. 1 2 1 2 3 3 4 4 5 5 6 6

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Table 5.

	Additions to the control wells							
	Controls:		Biotinylated	Anti-biotin antibody HRP-labelled				
	5109	130	9A4					
1	+	٠	-	+				
2	+	+	-	+				

Table 5. (continued)

	Additions to the control wells						
	Controls:		Biotinylated	Anti-biotin antibody HRP-labelled			
	5109	130	9A4				
3	+	-	+	+			
4	-	+	+	+			
5	-	+	-	+			
6	-	•	+	+			

[0089] The wells were washed three times with 200 µL/well of DPBS TW-20.

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[0090] Biotin conjugated monoclonal antibody 9A4 (Bt-9A4) was added to all peptide 130 (SEQ!D NO: 20) containing wells, all sample wells, and all control wells except 1,2, and 5. 100 µL/well of 9A4-Bt at 1 µg/mL in 0.1% NFDM DPBS was added to each well and the plate incubated for 40 min at 37° C.

[0091] Note: 9A4 was biotinylated using 37 μg of biotin-N-hydroxysuccinamide (Pierce Chemical) per mg of monoclonal antibody 9A4 for 2 hrs and then dialyzed over night using a 10 kD cut-off dialysis cassette (Pierce Chemical).

[0092] The wells were washed three times with 200 μL/well of DPBS TW-20.

[0093] Mouse monoclonal anti-biotin antibody conjugated with HRP (Jackson ImmunoResearch) was diluted 1/5000 in 0.1% NFDM DPBS and 100 µL/well was added to all wells and incubated for 30 minutes at RT.

[0094] The wells were washed three times with 200 μ L/well of DPBS TW-20.

[0095] One hundred microliters/well of 1-step Turbo (ready to use 3,3',5,5'-tetramethyl benzidine; Pierce Chemical) was added to each well and incubated at RT for approximately 10 minutes. Color development was stopped with 2N H₂SO₄. The results were read with a spectrophotometer at 450 nm.

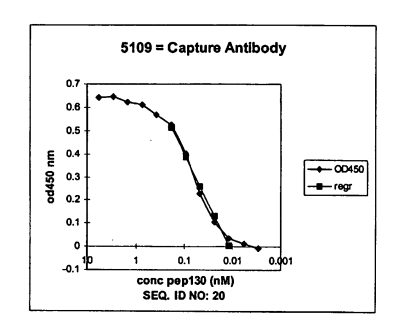
Table 6. Standard curve data for 5109 capture/Bt-9A4 detection sandwich assay

Cln7	pep130 (SEQ ID NO: 20)		log		lin	lin regi	ression
	ng/mL	nM	nM	OD450	regr	slope	intercept
1	10	5.88	0.769	0.64		0.42	0.70
2	5	2.94	0.468	0.65			
3	2.5	1.47	0.167	0.62			
4	1.25	0.735	-0.134	0.61			
5	0.625	0.368	-0.435	0.57			
6	0.313	0.184	-0.736	0.52	0.52		
7	0.156	0.092	-1.037	0.40	0.39		
8	0.078	0.046	-1.338	0.23	0.26		
9	0.039	0.023	-1.639	0.11	0.13		
10	0.020	0.011	-1.94	0.04	0		
11	0.010	0.006	-2.241	0.01			
12	0	0		-0.01			

[0096] From the concentrations of peptide 130 (SEQ ID NO: 20) and the resulting optical density reading at 450 nm, a standard curve was constructed. Again, other appropriate peptides or collagen fragments could be utilized to prepare a standard curve. The units needed were expressed in terms of equivalents of standard. In this case, the units were appropriate in terms of nM equivalents of peptide 130 (SEQ ID NO: 20).

[0097] Over the linear portion of the curve, a regression line was used to fit the data. In the given case, the standard curve was linear between 0.3125 nM and 0.0195 nM when the concentrations were given in the log scale (as in the example Figure) and the regression curve was obtained using just the linear portion of the curve. For samples that fall outside of the linear portion, the concentrations can be read off the graph or the samples may be diluted to fall within the standard portion of the curve, or the concentration of collagen fragments in the samples may be below the detection limit.

Figure 2. Standard curve for 5109 capture/Bt-9A4 detection sandwich assay.



[0098] The regression between log (nM) and OD450 nm gives a slope of 0.42 OD450/log(nM) and an intercept of 0.70 OD450 nm. When samples are run, the calibration curve can be used to determine the concentration of collagen fragment from the optical density of the sample.

Samples:

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[0099] In the given case, the unknown sample of human urine from an arthritic patient has an OD450 of 0.124

Log(Concentration) = (Sample OD450 - 0.70)/0.42 = - 1.36

Taking the anti-log, the concentration of fragment in urine = 0.44 nM

[0100] In another case, the standard curve gave a slope of 0.249 and an intercept of 0.638. An unknown sample of human osteoarthritis plasma had an OD450 nm of 0.172. The sample of osteoarthritic plasma had a concentration of 68 pM.

Example 5

Antibody 5109 can be used directly to measure the amounts of type II collagen fragment in a competition assay.

[0101] In an adaptation of concentration analysis ($BlAapplications\ Handbook$, Pharmacia Biosensor, June, 1994 Edition, p. 6-2 to 6-9), streptavidin (Pierce Chemical, Rockford, IL) at 100 µg/mL was conjugated with the Pharmada Amine Coupling Kit (Pharmacia Biosensor) to carboxylated dextran coated biosensor chips (Pharmacia Biosensor) at pH 4.0 using a flow rate of 5 µL/minute for 35 minutes. Typically, 2000 RU was added. Biotinylated peptide (100 ng/mL) having the sequence set forth in the Sequence Listing as SEQ ID NO: 19 was passed over the streptavidin chip at a flow rate of 5 µL/minute for 2 minutes; 144 RU of peptide was added to the streptavidin surface. MAb 5109 at a concentration of 6.3 µg/mL either alone or in mixtures with standard concentrations of peptide 054 (SEQ ID NO: 19) or mixtures of 5109 and dilutions of samples with unknown amounts of collagen fragments were passed over the peptide surface for 1 minute at a flow rate of 10 µL/min. The slopes of the linear portions of the association phase for each curve were analyzed with BlAevaluation version 2.1 software. A standard curve of competing peptide 054 (SEQ ID NO: 19) vs slope was constructed. The amount of collagen epitope in the samples was determined by comparison of a sample's slope to the slopes of the standard curve to calculate the amount of epitope. Between each injection,

the chip was cleaned with 0.1 N HCI for 30 seconds to remove antibody.

Table 7. 5109 mixed with standard amounts of peptide 054 (SEQ ID NO: 19).

	Dil. of			linear
l _	pep054		r0	regression
	M	log M	slope	values
1	1.34E-06	-5.87	0.187	
2	6.71E-07	-6.17	0.208	
3	3.36E-07	-6.47	0.375	
4	1.68E-07	-6.78	0.729	2.16
5	8.39E-08	-7.08	12.7	10.09
6	4.19E-08	-7.38	17.1	18.03
7	2.10E-08	-7.68	25.7	25.96
8	1.05E-08	<i>-</i> 7.98	22.5	
9	5.24E-09	-8.28	26.6	
10	2.62E-09	-8.58	24.0	
11	1.31E-09	-8.88	26.6	
12	6.55E-10	-9.18	25.6	

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Linear regression slope = -26.36

y-intercept = -176

Figure 3.

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Standard curve:
Inhibition of 5109a binding by
pep054 (SEQ ID No: 19)

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16-10 0.00000001 0.00000001 0.0000001 0.00001 0.00001 0.00001

pep054 conc. M

Sample:

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[0102] Supernatant of a collagenase-3 MMP13 digest of bovine nasal cartilage. The sample was run over the BIAcore chip diluted 2-fold, 4-fold and 8-fold. The calculated amounts of collagen in terms of 054 peptide (SEQ ID NO: 19) were determined. The results are given in column 4 of Table 8. After multiplying by the titer, a molar concentration (M) of collagen can be determined in terms of the peptide 054 (SEQ ID NO:19) standard.

Table 8. The titer and concentrations of the unknown sample in terms of peptide 054 (SEQ ID NO: 19) concentration.

	titer	slope	M (054)	nM (054)
5109	alone	6.648	Background	
5109+	1:2	3.539	4E-08	80
5109 +	1:4	4.885	2E-08	80
5109 +	1:8	5.825	8E-09	64

[0103] The consistency of the results (last column) after correction for dilution is shown by the agreement of the values calculated from the three separate dilutions (next to last column) of the unknown samples. An average value of 75 nM type II collagen fragments is obtained for the bovine nasal cartilage supernatant.

Example 6

Preparation of genetically engineered antibodies related to 9A4.

[0104] The basis for generating engineered antibodies and their subsequent evaluation as biologically active or relevant molecules is the cloning, assembly configuration and characterization of the V_L and V_H domains of the parent antibody. We have determined the V_L and V_H structural sequences of the subject antibodies and the uniqueness of the particular V_L - V_H combination that forms the active binding site to the antigens described in this invention. Before cloning the 9A4 variable region genes, it was necessary to determine the protein sequence of portions of the variable domains of the parent 9A4 IgG1 antibody so that when the variable domains were cloned, it could be ascertained that the correct variable domains were indeed obtained and not other ones derived from the myeloma fusion partner or an inactive pseudogene from the B-cell used in generating the hybridoma. Culture supernatant containing 9A4 IgG1 was generated by growing the 9A4 hybridoma in roller bottles. Supernatants were adjusted to pH 7.5 with dibasic sodium phosphate and the salt concentration adjusted with 3 M sodium chloride to a final concentration of 150 mM. Filtered (0.2 μ) supernatant was passed through a 15 mL bed volume of Protein G (Pharmacia) at a flow rate of 20 mL/min. After further washing the column with 150 mM NaCl solution, the antibody was eluted with 100 mM glycine pH 3.1. The antibody was isotyped using anti-sera from the Mouse Immunoglobulin Isotyping Kit (Boehringer Mannheim, Indianapolis, IN) and found to be an IgG1 class murine antibody with a kappa light chain.

[0105] It has been observed that some isolated proteins are "blocked" at their amino terminus. By "blocked" is meant that the amino acid residue at the amino terminus of the polypeptide chain has been chemically modified in its structure post-translationally by cellular action or some spontaneous chemical change in such a way that the polypeptide chain is resistant to Edman degradation. The Edman degradation method is the chemical procedure routinely used over the past 40 years for determining the amino acid sequence of proteins. Use of the Edman degradation technique, normally automated in a laboratory instrument known as a sequenator or protein sequencer, is a standard procedure well known to those skilled in the art of protein biochemistry.

[0106] A common mode of blocking is conversion of an amino-terminal glutaminyl residue into a pyroglutamyl residue. This occurs by cyclization of the glutaminyl residue to form a structure which is inaccessible to the Edman reaction because a new amide linkage is formed between the former alpha-amino group and the delta-carboxyl group. In such circumstances, the ability to obtain sequence information from the amino terminus of the protein depends on removal of the pyroglutamyl residue by a chemical or enzymatic method. An important method is to use an enzyme called pyroglutamate aminopeptidase (EC 3.4.19.3) to remove the pyroglutamyl residue. The blocked protein, which may be in solution or electroblotted to a membrane material such as PVDF (polyvinylidene difluoride), is treated with a solution of pyroglutamate aminopeptidase until a sufficient amount of the protein has been unblocked to allow successful determination of the amino acid sequence by automated Edman chemistry. Example methods for this are described in: Fowler EF, Moyer M, Krishna RG, Chin CCQ and Wold F, (1995) "Removal of N-Terminal Blocking Groups from Proteins" in Current Protocols in Protein Science., pp.11.7.1-11.7.17 (eds. Coligan JE, Dunn BM, Ploegh HL, Speicher DW., & Wingfield PT.), John Wiley, New York.

[0107] In the present work, the heavy and light chains of MAb 9A4 were separated by SDS-polyacrylamide gel electrophoresis with the use of a reducing agent (beta-mercaptoethanol) in the sample buffer. Following electrophoresis, polypeptides in the gel were electroblotted to a PVDF membrane and detected by staining with Coomassie Brilliant Blue R-250. Bands containing the heavy and light chains of 9A4 were then excised from the blot and separately treated with pyroglutamate aminopeptidase. In each case, it proved possible subsequent to this treatment to obtain aminoterminal sequence information by Edman degradation. Sequencing was performed on a Perkin-Elmer Applied Biosystems Model 494 Procise protein sequencer.

[0108] When it is desired to obtain internal (i.e. not N-terminal) amino acid sequence information from the protein, blotted samples of the protein may be digested with trypsin and the resulting digest fractionated by HPLC to afford individual peptides which may then be sequenced.

Specifics of the work were as follows:

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N-terminal de-blocking with pyroglutamate aminopeptidase (PGAP)

[0109] Antibody (9A4) was separated into its constituent heavy and light chains by SDS-PAGE on a Tris-Gly 4-20% polyacrylamide gel (Novex, San Diego). It was then electroblotted onto ProBlott (Perkin-Elmer Applied Biosystems, Foster City, CA) and the bands were visualized by Ponceau S (Sigma) staining.

[0110] The membranes containing 9A4 light chain were excised and incubated for 30 min in a buffer containing: 0.1 M sodium phosphate, 10 mM Na₂EDTA, 5 mM dithioerythritol, 5% glycerol, and 0.1% reduced Triton X-100. Pyroglutamate aminopeptidase (20 mg)(Boehringer Mannheim, Indianapolis, IN) was added to the vial, the contents mixed gently, and the reaction was incubated overnight at 37°C. The membranes were removed and washed extensively in water to remove all salts, detergent and enzyme. The membranes were then placed in the Applied Biosystems Model 494 protein sequencer (Perkin-Elmer) for N-terminal sequence analysis according to the directions provided by the manufacturer.

[0111] The heavy chain was treated in the same manner as above for the light chain.

[0112] The following N-terminal sequence results were obtained:

Sample	Sequence data file	Sequence
Light chain	9A4L_7 3-22-96	IVLTQSPVFMSASPGEKVTM (Note 1)
Heavy chain	9A4H_4 3-26-96	IQLVQSGPELKKPGQTVKI(S) (Note 2)

Residues in parenthesis indicate tentative calls.

Note 1: This sequence corresponds to residues 9 to 28 of SEQ ID NO: 33. Note 2: This sequence corresponds to residues 13 to 32 of SEQ ID NO: 32.

[0113] In addition to the N-terminal sequence data obtained above,internal sequencing of the 9A4 antibody light chain was also performed, according to the method developed from:

Femandez J, Andrews L, Mische SM, Anal Biochem 1994; 218: 112-117.

[0114] Four bands corresponding to the 9A4 light chain were excised from the ProBlott and incubated for 30 min in a buffer containing 10% acetonitrile, and 0.1 % reduced Triton X-100 in 0.1 M Tris-HCl pH 8.8. Sequencing Grade Modified Trypsin (0.2 mg)(Promega, Madison, WI) was added and the bands were incubated overnight at 37° C. The resulting peptides were extracted from the membrane by washing with 60% acetonitrile, 0.1% TFA, H₂O and sonication.

The peptides were separated by RP-HPLC on a Vydac C18 218TP column (1.0 x 250 cm)(Vydac, Hesperia, CA). The peaks were collected visually by hand and selected peaks were analyzed by automated Edman sequencing on a Model 494 protein sequencer as above.

[0115] The following N-terminal sequence results for the peptide fragments were obtained:

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Sequence data file	Sequence	
9A4L_10	DSTYSMSSTL (C, se	equence)
9A4L_12	LLIHATSNLASGVPV	
9A4L_13	FSGGGSGTSYSLTIS	SR (Note 2)
9A4L_14	XFNR	(C _r sequence)
9A4L_15	(H)NSYTCEATHK	(C, sequence)
9A4L_16	(Q)NGVLNGTSY	(C _s sequence)
9A4L 17	LEIIR	(Note 3)

Residues in parenthesis indicate tentative calls.

- Note 1: This sequence corresponds to residues 52 to 67 of SEQ ID NO: 33.
- Note 2: This sequence corresponds to residues 68 to 83 of SEQ ID NO: 33.
- Note 3: This sequence corresponds to residues 110 to 114 of SEQ ID NO: 33.

[0116] Mature 9A4 Light Chain (V_L-C_{kappa}) Amino Acid Sequence (comparison of sequence derived from DNA sequencing of the cloned V_L (see below) and the murine C_{kappa} (obtained from Kabat EA, Wutt, Perry HM, Bottesman KS and Foeller C.-"Sequences of Proteins of Immunological Interest, U.S. Department of Health and Human Services, NIH, 5th Edition, publication # 91-3242, 1991) with sequences obtained by Edman degradation)

- 1 QIVLTQSPVF MSASPGEKVT MTCSASSSVS YMYWYQQKPG SSPRLLIHAT SNLASGVPVR
- 61 FSGGGSGTSY SLTISRMEAE DAATYYCQQW RSYTRTFGGG TKLEII*RADA APTVSIFPPS
- 121 SEQLTSGGAS VVCFLNNFYP KDINVKWKID GSERQNGVLN SWTDQDSKDS TYSMSSTLTL
- 181 TKDEYERHNS YTCEATHKTS TSPIVKSFNR NEC

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Underlined amino acids have been sequenced by automated Edman sequencing [Also see table above]. The sequence obtained from the cloned DNA, when compared with the amino acid sequences from the fragments of the original antibody protein shown above, indicates that the gene cloned is the correct one for the 9A4 $V_{\rm L}$.

*Amino acid 106 above is usually a Lys residue in the J1 germline segment, but has been mutated to the lie residue shown above for the 9A4 V_L . Amino acid 106 marks the end of the V_L domain while amino acid 107 (Arg) marks the beginning of the C kappa domain.

Cloning and Determination of the V_L and V_H Sequences of MAb 9A4

[0117] The 9A4 hybridoma cell line was grown in HT media (Sigma) with 5% fetal calf serum (Hyclone). mRNA was extracted from a cell pellet containing 1 x 10^7 cells using the Pharmada Quick Prep mRNA Extraction Kit (Pharmacia) as per the manufacturer's protocol. cDNA was then synthesized for both the V_L and V_H gene segments using Boehringer Mannheim's cDNA Kit (Indianapolis, IN) as per the protocol provided by the manufacturer. The oligo used in the V_L cDNA reaction (named MLK) and set forth in the Sequence Listing as SEQ ID NO: 21 was specific for the murine light chain kappa region, while the oligo used in the V_H cDNA reaction was specific for a segment in the C_H2 domain of the murine heavy chain gamma region was named MHG. The sequence of MHG is set forth in the Sequence Listing as SEQ ID NO: 22. This oligo was designed to anneal to all 4 murine IgG isotypes, including IgG1, IgG2a, IgG2b and IgG3. There is a single base mismatch for IgG1 at nucleotide 5 of SEQ ID NO: 22, but it should be apparent to those skilled in the art that this would not prevent annealing and subsequent generation of cDNA.

[0118] Note: Oligos were synthesized by Oligos Etc (Wilsonville, OR), Genosys (The Woodlands, TX) or Perkin-Elmer Applied Biosystems (Foster City, CA).

[0119] The amino terminal amino acid sequence data was used to generate a possible oligo to isolate the correct

9A4 V_H gene by PCR as follows. Based on the 20 amino acid sequence presented above for the V_H amino terminal region, which corresponds to residues 13 to 32 of SEQ ID NO: 32, and assuming that the amino terminal amino acid of the mature form of the secreted antibody heavy chain was indeed a GIn residue [residue 12 of SEQ. ID. NO. 32], sequences of known antibodies were compared to that of 9A4 V_H using the Kabat data base: Sequences of Proteins of Immunological Interest, Volume II, 1991. Antibody V_H domains that matched the first 20 amino acids of the mature 9A4 V_H included: MAb 264 (Nottenburg C, St. John T, and Weissman IL J Immunol 1987;139: 1718-1726); MAbRFT2 (Heinrich G, Gram H, Hocher HP, Schreier MH, Ryffel B, Akbar A, Amlot PL, and Janossy G, J Immunol 1989; 143: 3589-3597; and MAb2H1 (Li Y-W, Lawrie DK, Thammana P, Moore GP and Shearman C W, Mol Immunol 1990; 27: 303-311). Since it is important to obtain the original DNA sequence corresponding to the amino terminus of the mature antibody, the 5' PCR oligo should be designed to anneal 5' (upstream) from the amino terminus, i.e. in the signal peptide segment. The DNA sequences of all the above 3 antibodies, with which the first 20 amino acids of the 9A4 V_H was identical, had known signal peptide DNA and amino acid sequences.

[0120] The DNA sequences are shown here: (Underlined nucleotides indicate differences between the sequences).

264: 5'- GG CTG TGG AAC TTG CTA TTC (SEQ ID NO: 23)

RFT2: 5' - GG GTG TGG ACC TTG CCA TTC (SEQ ID NO: 24)

2H1: 5'- GG GTG TGG ACC TTG CTA TTC (SEQ ID NO: 25)

[0121] These sequences code for amino acids -11 to -17 in the signal peptides for the V_Hs of the corresponding antibodies [Kabat]. The 5' oligo, MHMISC was thus designed to isolate the genuine 9A4 V_H. The sequence of MHMISC is set forth in the Sequence Listing as SEQ ID NO: 26.

[0122] For isolating the 9A4 V_L, a series of five degenerate oligo sets were designed to anneal to the leader peptide segments of known murine kappa light chains. These oligos were employed in 5 separate PCR amplifications. (See below). The sequences of the 5 degenerate oligo sets were provided by The Dow Chemical Company (Midland MI) and are hereby acknowledged. The 5' oligo set that gave a bona fide 9A4 V_L was MK3, the sequence of which is set forth in the Sequence Listing as SEQ ID NO: 27. These oligos code for residues -8 to -14 in the signal peptide.

[0123] The reverse primers for the V_L and V_H PCRs were designed from known sequences of the murine IgG1 heavy chain constant region (from the C_H 1 domain), called MIGG1CH1, the sequence of which is set forth in the Sequence Listing as SEQ ID NO: 28, and the constant region of the kappa light chain, called MLKN, the sequence of which is set forth in the Sequence Listing as SEQ ID NO: 29. These oligos were nested (upstream) relative to the 3' primers used in the cDNA synthesis.

[0124] The annealing segment begins at nucleotide 10 of SEQ ID NO: 29.

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[0125] The 9A4 V_H and V_L genes were amplified by PCR using the corresponding oligos described above and 0.41 μg of 9A4 hybridoma cDNA. The PCR was set up using a GeneAmp® Kit with Native Taq Polymerase (Perkin Elmer) and used according to the manufacturer's specifications. The temperatures of denaturation, annealing and polymerization were 94°C, 55°C, and 72°C for 45 s, 45 s, and 60 s seconds, respectively for the V_H PCR and for the V_L PCR the annealing temperature was changed to 60°C. The PCR was carried out for 36 cycles plus a final polymerization cycle of 72°C for 7 min followed by a 4°C hold. The PCR products were cloned in the pCR2.1 vector (Invitrogen, Carlsbad, CA) and also sequenced to determine and verify the DNA and derived amino acid sequences of the V_L and V_H . The oligos used for sequence determination were MK3 and MLKN for the V_L and MiGG1CH1 for the V_H and are set forth in the Sequence Listing as SEQ ID NOS: 27, 29 and 28, respectively.

[0126] The DNA sequences corresponding to 9A4 V_H and V_L are set forth in the Sequence Listing as SEQ ID NOS: 5 and 6, respectively. The derived amino acid sequences of the 9A4 V_H and V_L genes are set forth separately in the Sequence Listing as SEQ ID NOS: 32 and 33, respectively.

[0127] Surprisingly, the oligo MHMISC (SEQ ID NO: 26) successfully provided a VH sequence which corresponded exactly to the first 20 amino acids of the mature protein V_H determined by protein sequencing of 9A4 IgG1. Immediately 3' of the 5' PCR oligo, MHMISC (SEQ ID NO: 26), the DNA sequence in the leader peptide segment of the 9A4 V_H is almost identical to the corresponding segment of MAb 2H1, and is likely therefore to be derived from the same germline V_H gene as 2H1. There are only 3 differences in amino acid sequence between the 2 antibodies in the V_H regions; 1 in CDR1, 1 in CDR2 and 1 in FR3. These are likely somatic mutations that have occurred during the affinity maturation process for these two antibodies and may play a role in defining the specificity and affinity of each of the antibodies for their respective targets. The 9A4 V_H utilizes the murine $J_H 2$ joining segment gene and codes for residues 114 (within

the CDR3) to 126 of SEQ ID NO: 32.

[0128] The 9A4 variable heavy chain belongs to Kabat's Mouse Ig heavy chain Family II. (The V_H genes here identified were taken from the Kabat Database at http://immuno.bme.nwu.edu/famgroup.html). The 9A4 V_H heavy chain (SEQ ID NO: 5) most closely resembles Database ID number 001246. There are only 2 differences in these 2 V_H genes, one occurring in the FR1 (silent mutation) and the other in the CDR1, at amino acid position 45 (SEQ ID NO: 32) where 9A4 has an Ite and 001246 has a Met. It is most likely that both of these genes are also derived from the same germline V_H . [0129] We claim V_H genes in other antibodies related to the 9A4 V_H germline gene, such that when the V_H gene product forms a productive antibody $V_L - V_H$ pair with the V_L of this other antibody, it binds in a manner (specificity and affinity) analogous to 9A4.

- [0130] The 9A4 V_L gene and derived amino acid sequences are set forth in the Sequence Listing as SEQ ID NOS: 6 and 33, respectively. The derived amino acid sequence of the 9A4 V_L gene matches all the peptide fragments from protein sequencing of genuine 9A4 light chain as presented above. The 9A4 variable light chain belongs to Kabat's Mouse Kappa Family XI and is most similar to Database ID Number 006306. There are 10 nucleotide mismatches, resulting in 7 amino acid differences. Two of these differences occur in FR1, 2 in CDR2, 1 in FR3 and 2 in CDR3. Since the majority of the changes occur in the CDRs, it is most likely that these 2 genes are related by being derived from the same or at least very similar germline V_L.
 - We claim V_L genes in other antibodies derived or related to the 9A4 V_L germline gene, such that when the V_L gene product forms a productive antibody V_L - V_H pair with the V_H of this other antibody, it binds in a manner (specificity and affinity) analogous to 9A4.
- We also claim antibodies where both the V_L and V_H are derived from the 9A4 V_L and V_H germline genes, disclosed herein, such that when the V_L and V_H gene products form a productive V_L-V_H pair, this other or different antibody from 9A4 essentially binds in a manner (specificity and affinity) analogous to 9A4.
- We now have the basis from which to design and construct genetically engineered versions of the subject invention 9A4 antibody. Two examples are presented below, both single chain antibodies (scFv). In one case, the design is V_H-Linker-V_L-HIS-MYC tags in the vector pCANTAB6 and in the other the scFv is constructed in the opposite orientation with a different linker, that is, V_L-Linker-V_H-FLAG tag, in the vector pATDFLAG. These examples are not meant to be limiting, but to those skilled in the art, it will readily be apparent that the newly characterized V_H and V_L domains of 9A4 (SEQ ID NOS: 5 and 6) can be utilized in part or in whole to obtain previously described antibody types, such as Fab's, or novel configurations that utilize only some critical portion of the V domains.

Construction of 9A4 scFv (VH-linker-VL) in pCANTAB6

- [0131] SOEing (Splicing by Overlap Extension) PCR primers were utilized to prepare for assembly of the V_H and V_L fragments into a scFv antibody.
- [0132] Two primers were designed and obtained from Perkin Elmer for the V_H region: for the 5' V_H end primer an Sfi I site was added while at the 3' end for the V_H primer, an overlapping sequence with the (Gly₄Ser)₃ linker was added. The 5' V_H primer was called 9A4VH5CAN and the 3' V_H primer was called 9A4H3CAN, which correspond to SEQ ID NOS: 34 and 35 in the Sequence Listing.
 - [0133] For the V_L, a 5' end primer with overlap into the Gly₄Ser linker region and a 3' primer incorporating a Not I restriction site were designed and also obtained from Perkin Elmer. The 5' primer was called 9A4VL5CAN and the 3'V_L primer was called 9A4VL3CANILE, which correspond to SEQ ID NOS: 36 and 37 in the Sequence Listing.
 - **[0134]** The V_H and V_L DNA components were amplified by PCR and joined subsequently by an assembly, pull-through SOE-PCR step. Following the SOE-PCR reaction, a band in an agarose gel, which was identified to be ~700 bp, was excised and gel purified using the QIAquick Gel Purification Kit (QIAgen) as per the manufacturer's directions.
- The resulting DNA was digested with Sfi I and Not I and used in a ligation with the pCANTAB6 vector DNA (obtained from Cambridge Antibody Technology, Melboume, Cambridgeshire, UK) which was also digested with the same restriction enzymes. The ligated DNA was then used to transform competent E. coli TG1 cells by electroporation. The basic protocol for generating the competent E.coli TG1 cells was as follows:
 - [0135] 2YT media (Bio101, LaJolla, CA)500 mL prewarmed to 37°C in a 2 L conical flask was inoculated with 2.5 mL of fresh ovemight culture of TG1 cells. They were grown with vigorous aeration (300 rpm) at 37°C until the OD at 600 nm was 0.2 to 0.25, usually 1-1.5 h. later. The flasks were chilled for 30 min on ice, then poured into 250 mL centrifuge bottles and spun at 4,000 rpm for 15 min in a prechilled (4°C) Sorvall centrifuge to pellet the cells. The cells were resuspended in the original volume of prechilled water, and spun again as above to pellet the cells.
 - [0136] The cells were resuspended in half the original volume, i.e. 250 mL of prechilled water and left on ice for 3 min, were resuspended and spun again as above.
 - [0137] The cells were then resuspended in 20 mL of prechilled 10% glycerol, transferred to a prechilled 50 mL Falcon tube, left on ice for 15 min, and centrifuged at 3,500 rpm, at 4°C for 10 min in a benchtop centrifuge.
 - [0138] The cells were finally resuspended in 1.0-2.5 mL prechilled 10% glycerol and used directly in the electropo-

ration step.

[0139] Ligated DNA (25-250 ng pCANTAB6-Sfi I/Not I vector with 9A4 V_H-L-V_L-Sfi I/Not I) was electroporated into the competent *E. coli* TG1 cells as follows:

[0140] The ligated DNA was ethanol precipitated per standard protocol. The DNA pellet was dissolved in 10 μ L of sterile deionized distilled water.

[0141] The DNA (up to 10% of the total cell volume) was added to 100 μL of cells and transferred to a prechilled electroporation cuvette (Biorad) and left on ice.

[0142] The electroporation Gene Pulser apparatus (Biorad) parameters were set to 25 µFD, 2.5 kV and the pulse controller set to 200 ohms. The cuvette was dried with a tissue, placed in the electroporation chamber and pulsed once. Time constants in the 3.5-4.8 msec range were typically obtained. The electroporated cells were immediately diluted with 1 mL of 2YT medium supplemented with 2% glucose. The cells were transferred to a 15 mL Falcon tube and shook at 37°C for 1 h.

[0143] The transformed cell mixture (20-1000 μL) was plated on appropriate size agar media plates containing 2YT, 2% Glucose and 100 ug/mL ampicillin (2YTAG).

[0144] Initially, a clone was obtained (p9A41CAT3-2) that had a 5 base insertion at the Not I site, which put the downstream sequence out-of-frame. In order to correct this, a new oligo was made called 9A4NOTFIX3, the sequence of which is set forth in the Sequence Listing as SEQ ID NO: 38.

[0145] A PCR amplification using *E. coli* cells containing p9A41CAT3-2 as the target for the annealing oligos was performed using the Advantage KlenTaq Polymerase Mix (Clontech, Palo Alto, CA) as per the manufacturer's protocol. The annealing oligos (35 pmol each) were 9A4NOTFIX3 and pUC19R. The sequence of pUC19R is set forth in the Sequence Listing as SEQ ID No. 39; it anneals upstream from the Sfi I site.

[0146] The PCR cycles (in a Perkin Elmer Cetus Model 9600 thermal cycler) were set up as follows: For the first cycle, denaturation of DNA was for 1 min at 94°C, annealing for 45 s at 60°C and polymerization for 1.5 min at 68°C. For cycles 2-31, the same temperatures and times were used, except the denaturation times were reduced to 30 s. For the final cycle (32) the polymerization was set up for 5 min, after which time the cycler cooled the sample to 4°C. The TA cloning system (Invitrogen) was used to clone the resulting PCR products in the plasmid pCR2.1 using the protocol suggested by the manufacturer. Twelve white colonies were screened for inserts using the oligos M13 Forward and M13 Reverse (Invitrogen) (as set forth in the Sequence Listing as SEQ. ID. NOS. 53 and 54, respectively) by PCR using KlenTaq Polymerase as above. Three colonies produced inserts of the correct size on agarose gel electrophoresis. One of these with the correct DNA sequence for the 9A4 V_H-Linker-V_L construct was chosen for further work. The DNA insert containing the scFv gene was obtained by digestion of the pCR2.1 derivative with Sfi I and Not I, purified using the QIAquick Gel Extraction kit and ligated with the pCANTAB6 vector digested with the same restriction enzymes. The DNA was electroporated into competent E. coli TG1 cells as described above and resulted in a clone named p9A4ICAT7-1 (ATCC 98593) which contained an active scFv. The DNA sequence of this 9A4 scFv is set forth in the Sequence Listing as SEQ ID NO: 7, while the amino acid sequence is set forth separately as SEQ ID NO: 40. Note that the GTG codon, beginning at position 29 of SEQ ID NO: 7 is the start codon (Met). The sequencing oligos used were pUC19R and FDTETSEQ which are set forth in the Sequence Listing as SEQ ID NOS: 39 and 41 respectively. [0147] See the information for SEQ ID NOS: 7 and 40 in the Sequence Listing for specific features of the DNA and amino acid sequences for this scFv antibody.

[0148] The genetically engineered 9A4 V_H-Linker-V_L format scFv antibody was expressed in *E. coli* TG1 cells, and the antibody was purified using NTA-Ni agarose (QlAgen) affinity chromatography, followed by Superdex 75 gel filtration chromatography to isolate monomer scFv species. The binding to the parental antigen of the scFv was evaluated on the BlAcore (see below). *E. coli* containing the plasmid pA4lCAT7-1 has been deposited with the American Type Culture Collection as ATCC 98593.

Construction of 9A4 scFv (V_L-linker-V_H) in pATDFLAG

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[0149] SOEing oligos were also prepared for PCR-SOE assembly of the V_H and V_L fragments in the opposite configuration, i.e.V_L-linker-V_H, relative to the example presented above in p9A4ICAT7-1. Two primers were designed for the V_L - linker region: at the 5' end an Nco I site was added and at the 3' end, an overlapping sequence with the 25 amino acid linker sequence set forth in the Sequence Listing as SEQ ID NO: 42 (Pantoliano MW, Bird RE, Johnson S, Asel ED, Dodd SW, Wood JF, and Hardman KD Biochemistry 1991; 30: 10117-10125) was added. The 5' V_L primer was named 9A4VL5ATD and the 3'V_L primer was named 9A4VL3ATDILE. The sequences of these oligos are set forth in the Sequence Listing as SEQ ID NOS: 43 and 44 respectively.

[0150] For the V_H, a 5' end primer with overlap into the linker region and a 3' end PCR primer with an added Nhe I site was designed. The 5' V_H primer was named 9A4VH5ATD and the 3' V_H primer was named 9A4VH3ATD. The sequences of these oligos are set forth in the Sequence Listing as SEQ ID NOS: 45 and 46 respectively.

[0151] The V_L and V_H components were amplified by PCR. The V_L and V_H were then joined in the linker region, by

a SOE-PCR using the oligos 9A4VL5ATD (SEQ ID NO: 44) and 9A4VH3ATD (SEQ ID NO: 46). DNA at the correct size, ca. 700 bp was excised out from a 1% agarose gel and eluted using the QlAquick gel elution kit. The resulting DNA was trimmed at the ends with the restriction enzymes Nco I at the 5' end and Nhe I at the 3' end. This was ligated with the expression vector pATDFLAG (PCT WO 93/12231) treated with the same restriction enzymes. Competent *E. coli* DH5α cells were transformed with the ligation as per the manufacturer's protocol and plated on agar plates containing 20 μg/mL of chloramphenicol as the selective agent. Two of the clones that were sequenced, p9A4IF-5 and p9A4IF-69, had no PCR or construction errors. The sequencing oligos were: UNIVLSEQ-5' (SEQ ID NO: 30) and TERMSEQ(-) (SEQ ID NO: 31).

[0152] E. coli containing p9A4IF-5 was chosen for further work and expression of the engineered 9A4 scFv antibody. The DNA sequence of this scFv is set forth in the Sequence Listing as SEQ ID NO: 8, while the derived amino acid sequence is set forth separately as SEQ ID NO: 47. Specific features of the V_L-L-V_H-FLAG 9A4 scFv, such as signal peptide, linker and tag locations are indicated in the Sequence Listing for SEQ ID NOS: 8 and 47. It will be obvious to those skilled in the art that the engineered antibody described could be expressed not just from E. coli, but from other organisms as well, including, but not limited to-P. pastoris, Baculovirus, Bacillus species, mammalian cells etc.

[0153] For expression and purification of this scFv product from *E. coli*, 1-2 L cultures of LB broth containing 20 µg/mL of chloramphenicol were grown ovemight at 37°C. Cells were pelleted in a Sorvall centrifuge using a GS-3 rotor. In preparation for affinity chromatography using an M2 affinity column (Kodak, New Haven, CT) (which is specific for the FLAG epitope shown in the sequence above) the pelleted *E. coli* cells were processed either in a Tris/EDTA/sucrose media to isolate the periplasmic fraction or were sonicated (Soniprep sonicator) directly in a minimal volume of DPBS buffer. The affinity column was washed extensively with DPBS to remove any unbound materials after loading the crude scFv sample. The scFv antibody was eluted using 0.1 M glycine-HCl pH 3.1. The monomer scFv species was isolated by Superdex-75 gel filtration chromatography (Pharmacia). The antibody was judged to be homogeneous by SDS-PAGE and staining with Coomassie Brilliant Blue R-250. Antibody was quantitated spectrophotometrically at OD 280 nm, where an absorbance of 1.4 was defined to be equivalent to 1.0 mg/mL scFv, using a 1.0 cm pathlength quartz cuvette. *E. coli* containing the plasmid p9A4IF-5 was deposited with the American Type Culture Collection as ATCC-98592.

[0154] Binding to antigen was evaluated on the BIAcore for both the p9A4ICAT7-1 and p9A4IF-5 scFv purified gene products as follows. A streptavidin chip was loaded with biotinylated peptide of SEQ ID NO: 14 as given in Example 1 above. Both scFv constructs were shown to bind antigen, i.e., compared to the parent 9A4 IgG which binds with a K of 1.2x10-7 M, the 9A4IF-5 scFv has a K of 1.1 x10-7 M. For the 9A4ICAT7-1 scFv, the off-rate was 1.2X10-3 sec-1 compared to 1.76x10-2 sec-1 for 9A4 IgG (parent antibody). These data indicate that the affinity of the engineered antibodies were at least as good or better than the parent.

[0155] As evidenced by the specificity and kinetic data determined by BIAcore, the 2 engineered antibodies containing the 9A4 V_L and V_H domains described above have the desired characteristics for using them in the quantitative measurement assays described in Examples 3 and 4 above. Other engineered antibodies comprised of some or all of the 9A4 V_L and V_H disclosed herein having the affinity and specificity of the parent 9A4 antibody, would therefore be considered to be in the scope of claims for this invention.

Example 7

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Preparation of the genetically engineered antibodies related to 5109.

[0156] Before cloning the 5109 variable region genes, it was necessary to determine the protein sequence of portions of the variable domains of the parent 5109 antibody so that when the variable domains were cloned, it could be ascertained that the correct variable domains were indeed obtained and not other ones derived from the myeloma fusion partner or an inactive pseudogene from the B cell used in generating the hybridoma. Culture supermatant containing 5109 was generated by growing the 5109 hybridoma in roller bottles. Supermatants were adjusted to pH 7.5 with dibasic sodium phosphate and the salt concentration adjusted with 3 M sodium chloride to a final concentration of 150 mM. Filtered (0.2 µ) supermatant was passed through a 15 mL bed volume of Protein G (Pharmacia) at a flow rate of 20 mUmin. After further washing the column with 150 mM NaCl solution, the antibody was eluted with 100 mM glycine pH 3.1. The antibody was isotyped using anti-sera from the Mouse Immunoglobulin Isotyping Kit (Boehringer Mannheim) and found to be an IgG1 class murine antibody with a kappa light chain constant domain.

[0157] In the present work, the heavy and light chains of MAb 5109 were separated by SDS-PAGE with the use of a reducing agent (beta-mercaptoethanol) in the sample buffer. Following electrophoresis, polypeptides in the gel were electroblotted to a PVDF membrane and detected by staining with Coomassie Brilliant Blue R-250. Bands containing the heavy and light chains of 5109 were then excised and subjected to Edman degradation. Sequencing was performed on a Perkin-Elmer Applied Biosystems Model 494 Procise protein sequencer as per the manufacturer's protocols. The sequences of the heavy and light chains that were obtained are shown in Table 9, and correspond to residues 1 to 40

of SEQ ID NO: 48 for the VH and to residues 1 to 39 of SEQ ID NO: 49 for the VL.

Table 9. Results of N-terminal amino acid sequence of 5109.

	Heavy	1	Light
Res#	chain	Res#	chain
1	E	1	D
2	V	2	V
3	Q	3	V
4	L	4	М
5	V	5	T
6	E	6	Q
7	S	7	T
8	G	8	Р
9	G	9	L
10	G	10	T
11	S	11	L
12	V	12	S
13	Q	13	V
14	Р	14	T
15	G	15	i i
16	G	16	G
17	S	17	Q
18	L	18	S
19	K	19	Α
20	L	20	S
21	S	21	1
22		22	
23	Α	23	
24	Α	24	K
25	S	25	S
26	G	26	S
27	F	27	Q
. 28	Т	28	
29	F	29	L
30	N	30	L
31	T	31	G
32	Y	32	(S)
33	G	33	D
34	M	34	(G)
35	S	35	L
36	W	36	Т
37		37	Y
38	R	38	(L)
39	Q	39	
40	T	40	

Tentative amino acids are indicated by (). Note: Position 22 in the V_{H} and position 23 of the V_{L} is normally Cys, and cannot be determined by Edman degradation.

Cloning and Determination of the V_L and V_H Sequences of MAb 5109

[0158] The 5109 hybridoma cell line was grown in HT media (Sigma) with 5% fetal calf serum (Hyclone). Cells were pelleted (2.5 x10⁷ cells / pellet) and frozen at -80°C until use. Extraction of mRNA (Oligotex™ Direct mRNA Kit; QIA-GEN) was carried out according to the manufacturer's directions. cDNA was then synthesized for the V_L and VH regions using Boehringer Mannheim's First-Strand cDNA Synthesis Kit. The oligo used in the V_L cDNA reaction was specific for the murine light chain kappa region (MLK) while the oligo used in the V_H cDNA reaction, MHG, was specific for a segment in the murine heavy chain C_H2 gamma region. The sequences of oligos MLK and MHG are set forth in the

Sequence Listing as SEQ ID NOS: 21 and 22, respectively.

[0159] PCR primers were designed for the N-terminal sequence of the mature, secreted forms of the heavy and light chains based on the amino acid sequences that were obtained for the V_L and V_H by Edman degradation. The sequences were compared with the Kabat database; the 5109 V_H was found to be most similar to members of Kabat subgroup IIID, while the 5109 V_L was most similar to members of Kabat subgroup II.

[0160] The sequences of the 5' V_H primer and 5' V_L primer were named 51-09V_H5' NDe and 51-09V_L5' NDe respectively, and are set forth in the Sequence Listing as SEQ ID NOS: 50 and 51.

[0161] Reverse primers were designed from known sequences of the IgG1 heavy chain constant region (to a segment in the C_H1 domain) and the constant region of the kappa light chain. Both of these 3' oligos are 5' (upstream) of the original oligos used to generate the cDNA. The 3' V_H primer, named MIGG1CH1 and the 3' V_L primer named MULK2 are set forth in the Sequence Listing as SEQ ID NOS: 28 and 52, respectively.

[0162] The resulting PCR products for the V_L and V_H were ligated into pCR2.1 and representative clones were chosen for subsequent DNA sequencing. DNA sequencing was performed on an Applied Biosystems Model 373 Stretch Sequencer and was set up and operated according to the protocols provided by the vendor. For DNA sequence determination, the Invitrogen commercial sequencing oligos M13F and M13R were used, the sequences of which are set forth in the Sequence Listing as SEQ ID NOS: 53 and 54.

[0163] The first 21 amino acids that were determined by Edman degradation for the 5109 V_L and V_H mature amino termini were found to be identical to the corresponding amino acid sequences derived from the DNA sequence of the corresponding genes that were cloned here by PCR.

[0164] The DNA sequences of the 5109 V_H and V_L domains are set forth in the Sequence Listing as SEQ ID NOS: 10 and 11, respectively. The amino acid sequences of the 5109 V_H and V_L are set forth separately in the Sequence Listing as SEQ ID NOS: 48 and 49, respectively. It should be noted that while the DNA, as presented, codes for the correct amino acids in the amino terminal segments of each of the V_L and V_H seqments corresponding to the PCR annealing oligos, the exact codons for the antibody amino acid segments corresponding to these oligos, as they would have occurred in the original hybridoma DNA are not unequivocal. The 5109 V_H utilized the J_H3 joining segment gene with 2 mutations in the codon that would otherwise code for a Thr residue (positions 328 and 330 of SEQ ID NO: 10), but in the 5109 V_H is an Ala residue (position 110 of SEQ ID NO: 48).

[0165] The 5109 variable heavy chain is most related to Kabat's Mouse Ig heavy chain Family XIV and most closely resembles Database ID number 002754. There are 24 nucleotide differences between these 2 V_H genes, resulting in 14 amino acid differences. It is most likely, based on this high number of differences, that these two genes are not derived from the same germline gene. It is still possible, however, that the 2 are derived from the same germline and that both have been heavily mutated in the *in vivo* affinity maturation process and the resulting divergence was amplified.

[0166] We claim V_H genes in other antibodies related to the 5109 V_H germline gene, such that when the V_H gene product forms a productive antibody V_L - V_H pair with the V_L of this other antibody, it binds in a manner (specificity and affinity) analogous to 5109.

[0167] The 5109 V_L utilized a J5 joining segment and codes for amino acids 102 to 112 of SEQ ID NO: 49. The CDR3 for this antibody is relatively rare because of the Cys-94 residue contained therein. The CDR3 extends from residues 94 to 102 of SEQ ID NO: 49. When 5109 single chain antibodies were engineered with a 5109 V_L (see below) 2 versions were made, one with the parental Cys-94 and another with Ser substituting the Cys-94.

[0168] The 5109 V_L belongs to Kabat's Mouse Kappa Family VI and is most similar to Database ID Numbers 005841, 005842, 005843, and 005844. The 4 antibodies in the database are identical in their nucleotide sequences, so a comparison with the 5109 V_L can be made to all 4 of them at the same time. There are 12 nucleotide mismatches, resulting in 8 amino acid differences. One of these differences occur in FR1, 3 in CDR1, 1 in FR2, 1 in CDR2, 1 in FR3 and 1 in CDR3. These changes occur throughout the V_L, but are focused on the CDRs, with 5 out of the 8 differences being in these hypervariable segments. Therefore it is most likely that these genes are related by being derived from the same or at least very similar germline V_I.

[0169] We claim V_L genes in other antibodies related to the 5109 V_L germline gene, such that when the V_L gene product forms a productive antibody V_L - V_H pair with the V_H of this other antibody, it binds in a manner (specificity and affinity) analogous to 5109.

[0170] We also claim antibodies where both the V_L and V_H are derived from the 5109 V_L and V_H germline genes, disclosed in this patent, such that when the V_L and V_H gene products form a productive V_L - V_H pair, this other or different antibody from 5109 essentially binds in a manner (specificity and affinity) analogous to 5109.

Construction of 5109 scFv engineered antibody (VH-Linker-VL) in pUC119

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[0171] SOEing (Splicing by Overlap Extension) PCR primers were utilized to prepare for assembly of the V_H and V_L components. All of the oligos utilized in 5109 scFv construction were synthesized in-house using a Beckman Oligo 1000M DNA synthesizer (Fullerton, CA). These five PCR primers, called 5109 VH 5', 5109 VH 3', 5109 VL 5', 5109 VL

3' SER and 5109 VL 3' CYS are set forth in the Sequence Listing as SEQ ID NOS: 55, 56, 57, 58 and 59 respectively. [0172] The oligo 5109 VL 3' Ser (SEQ ID NO: 58) was used to change Cys-94 of SEQ ID NO: 49 to Ser-94 in the scFv construct, (corresponds to the Cys residue at position 248 of SEQ ID NO: 63) to potentially improve the stability of the resulting scFv. Another primer, 5109 VL 3' Cys (SEQ ID NO: 59) was also synthesized in order to retain the original Cys sequence.

[0173] Following the SOEing reaction, DNA was amplified by PCR and the product sequenced directly for sequence confirmation.

[0174] DNA sequence was verified using an Applied Biosystems Model 373 Stretch Sequencing Unit as per the directions of the manufacturer. The following oligos were used for sequencing of potential 5109 scFv engineered antibodies ligated directly into pUC119: pUC19R, MycSeq10, Gly4Ser5', Gly4Ser3', the sequences of which are set forth in the Sequence Listing as SEQ ID NOS: 39, 60, 61 and 62, respectively.

[0175] For those 5109 DNA cassettes ligated into the pCR2.1 vector, the M13F and M13R oligos purchased from InVitrogen (SEQ ID NOS: 52 and 53) were utilized for sequence priming reactions, along with SEQ ID NOS: 61 and 62, as two internal oligos.

[0176] Clones directly ligated into pUC119 contained a large number of PCR errors. An acceptable clone (H55) was identified however, in the pCR2.1 vector. This construct was then subcloned by digesting the scFv with Sfi I and Not I and ligated in the pUC119 vector similarly digested. Clones were screened for insert by PCR amplification using pUC19R and MycSeq10 oligos (SEQ ID NOS: 39 and 60). Three clones were submitted for sequencing. One clone was identified with the desired sequence and is designated p5109CscFv7 (ATCC 98594); the DNA and derived amino acid sequences are set forth in the Sequence Listing as SEQ ID NOS: 9 and 63, respectively. Sequence features of this engineered antibody are given in SEQ ID NO: 63. In generating the 5109 scFv by PCR, a mutation (PCR error) occured at nucleotide 738 of SEQ ID NO: 9, changing the amino acid coded for by the altered codon from valine to alanine. This corresponds to the Ala residue at position 237 of SEQ ID NO: 63 (the 5109 scFv sequence). As a conservative difference, it was considered unimportant in terms of affecting the activity of genetically engineered products and was thus allowed to be carried forward in the V_L of the scFv constructs described. Of course, for those skilled in the art, it will be apparent that this PCR error could also be corrected and the original Val residue obtained at this position.

[0177] To verity that the new single chain constructs retained the binding properties of the parent molecule, the 5109 scFv was expressed in *E. coli* and purified.

[0178] A 2YT starter culture (50 mL) containing 100 ug/mL ampicillin and 2% glucose was inoculated with 50 uL of -80°C glycerol stock. Cultures were incubated ON at 30°C with shaking at 300 rpm. Each of six 2 L flasks containing 2YT media supplemented with 100 ug/mL ampicillin and 0.1% glucose were inoculated with 5 mL of the overnight starter culture. Cultures were incubated 30°C until turbid and subsequently induced by adding IPTG isopropyl B-D-thiogalactopyramoside (Boehringer Mannheim) to a final concentration of 1 mM. Cultures were grown for an additional 4 hr for production of scFv. Cells were centrifuged at 5000 x g for 10 min. Cell pellets were stored at -20°C until processed.

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[0179] For scFv purification, cell pellets were resuspended in TES (0.2 M Tris-HCI, 0.5 mM EDTA, 0.5 M sucrose). Following resuspension, a 1:5 dilution of the above TES buffer containing protease inhibitors (Complete Protease Inhibitor Cocktail, Boehringer Mannheim) was added. This preparation was allowed to incubate at 4°C for 30 min. Following incubation, cells were pelleted at 12,000xg for 15 min. MgCl₂ was added to the resulting supernatant to a final concentration of 5 mM. Ni-NTA agarose (QlAgen) was washed 1 X in PBS containing 300 mM NaCl, 15 mM imidazole, 0.2% Triton-X, pH 7.4. Washed agarose was then added and the slurry was allowed to incubate for 30 min at 4°C. Ni-NTA agarose beads + scFv were then washed 4X as previously described. The scFv was then eluted in wash buffer containing 250 mM imidazole. The eluate was desalted over a NAP-25 column (Pharmacia-Biotech, Uppsala, Sweden) and concentrated using a Centriprep concentrator device (Amicon, Beverly, MA). Products were electrophoresed on SDS-PAGE and visualized by silver staining. The resulting scFv products for p5109CscFv7 (Cys) and p5109SscFvA9 (Ser) were approximately 15% and 75% pure, respectively.

[0180] Using Origen methodology (Technical manual, Origen Instrument, Igen Corporation, Gaithersburg, MD), binding to biotinylated peptide 225 (having the sequence set forth in the Sequence Listing as SEQ ID NO: 64) by the 5109 scFv species was measured. Peptide 225 was prepared by Anaspec. Peptide 225 was added to 800 μg/mL streptavidin coated magnetic beads (Dynabeads, Igen, Gaithersburg, MD) to bring the final concentration to 10 nM and incubated for 15 minutes. The new genetically engineered scFv antibody was added to the peptide /bead solution for 30 min with shaking. The 9E10 anti-myc tag ruthenylated MAb was added in 200 μL Igen assay buffer (Igen, Gaithersburg, MD) and the ECL signal read on the Origen Instrument (Igen). MAb 9E10 was generated and purified from the 9E10 cell line, which was obtained from the ATCC. The MAb was ruthenylated using the N-hydroxysuccinamide derivative Origen TAG-NHS Ester (Igen) according to the manufacturer's instructions. In the absence of 5109 scFv, a background signal of 2995 ECL units was obtained. Addition of the 5109 scFv Cys construct (p5109CscFv7) resulted in 536,997 ECL units. Addition of the 5109 scFv Ser construct (p5109SscFvA9) resulted in 694,253 ECL units. These results demonstrate that both 5109 scFv genetically engineered antibodies were biologically active and bound to the same collagen-

related peptide fragment as MAb 5109 does. A culture of *E. coli* containing p5109CscFvA9 was deposited with the American Type Culture Collection as ATCC-98594.

[0181] The specificity data determined by the Origen technology demonstrates that the engineered antibody containing the 5109 V_L and V_H domains described above have the desired characteristics for using it in the quantitative measurement assays described in Examples 3 and 4 above. Other engineered antibodies comprised of some or all of the 5109 V_L and V_H disclosed here, having the affinity and specificity of the parent 5109 antibody would therefore be considered to be in the scope of claims for this invention.

[0182] It should also be noted that engineered antibodies comprised of a combination of the 9A4 and 5109 V_L and V_H domains, such as a bispecific scFv dimer of the composition: $5109V_L$ -Linker- $5109V_H$ -Linker- $9A4V_L$ -Linker- $9A4V_H$ -Tag(s) would also be useful, as the only antibody reagent in Examples 3 and 4 above. The difference would be that a single bispecific reagent could be used in a one step ELISA rather than using 9A4 and 5109 separately. To those skilled in the art, it is obvious that a number of genetic compositions comprising the subject antibodies' variable domains could be joined to make a variety of bispecific molecules and thus simplify the assays presented in Examples 3 and 4. The avidity of such molecules could be such that the overall sensitivity of the assay may also be significantly improved.

Example 8

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[0183] Mutations or differences in amino acid sequences of antibodies related to the subject invention antibodies can retain the binding properties (affinity and specificity) and therefore the utility of the parent antibody. This is demonstrated below by generating a series of mutants in the CDR3 of the 9A4 V_H. To those skilled in the art, it is apparent that other mutations in other regions of the same germline V_L and V_H genes of both 9A4 and 5109 can give antibodies of the same or of better binding properties relative to the original antibodies disclosed in this invention.

Generation of 9A4 VH CDR3 Region Mutants

[0184] The pCANTAB6 derivative of the 9A4 scFv, namely p9A4ICAT7-1 presented in Example 6 above, was used as the starting material to generate mutants in the CDR3 segment and the Vernier residues immediately adjacent to the CDR3. The parent DNA sequence of the CDR3 V_H region being targeted for mutation comprised nucleotides 383 to 409 of SEQ ID NO: 7. The derived amino acids corresponding to these nucleotides are residues 119 to 127 of SEQ ID NO: 40. The CDR3 begins at residue 121 and ends at 126. To introduce the random mutations in this area, the V_H and V_L regions were amplified separately by 2 PCRs. In the first PCR, oligos (obtained from Oligos Etc.) pUC19R (SEQ ID NO: 39) and 9A4MUT (set forth in the Sequence Listing as SEQ ID NO: 65) were used to amplify the V_H portion, where 9A4MUT was the oligo which introduced the mutations.

[0185] Nucleotides 25 to 51 in 9A4MUT (SEQ ID NO: 65) were 10 % spiked. In other words, the sequence as written accounted for 90 % of the nucleotide added at each position 25-51, while the other 3 nucleotides, in each position, 25 through 51, were introduced to the growing oligo chain at 3.3 % each. This was accomplished by methods well known in the art of oligonucleotide synthesis. The fact that random nucleotides were indeed introduced in this defined region will be discussed below. The second PCR utilized the 2 oligos: (also obtained from Oligos Etc) 9A4L5 (SEQ ID NO: 66) and FDTETSEQ (SEQ ID NO: 41). This produced the Linker-V_L portion up to the Not I site in p9A4ICAT7-1 at the 3' end and with overlap into the V_H at the 5' end that would allow subsequent annealing and assembly with the mutated V_H PCR products from the first PCR.

[0186] The PCR was set up as follows. For the mutant V_H, 50 pmol of each of the oligos pUC19R (SEQ ID NO: 39) and 9A4MUT (SEQ ID NO: 65) were used in 100 microliter reactions. The template DNA target was an aliquot of SNAP (Invitrogen) purified plasmid DNA-p9A4ICAT7-1. Taq Polymerase (Perkin Elmer) was used in the 30 cycle PCR. Denaturation, annealing and polymerase reaction times and temperatures were 94°C for 1 min, 55°C for 1 min and 72°C for 2 min, respectively.

[0187] For the 30th cycle the polymerization reaction was extended for a total of 10 min, before cooling the reaction to 4°C. For the V_L, which would overlap with the V_H species, PCRs were set up using both KlenTaq polymerase (Clontech) and Taq Polymerase (Perkin Elmer). DNA products were purified with the QlAgen gel extraction kit. The PCR assembly reaction for the mutated V_H and the V_L was performed using aliquots (1-2 microliters) of each of the purified PCR products in a 25 cycle total/2 temperature cycling: 94°C for 1 min, 65°C for 4 min and holding at 4°C at the end. No oligos were used for this step. For the PCR pull-through of the mutated 9A4 assemblies, 5 microliters of unpurified assembly reaction was used as the template and the oligos pUC19R (SEQ ID NO: 39) and FDTETSEQ (SEQ ID NO: 41) as the annealing primers. Correct size products were observed at ca. 900 bp and were gel purified using the QlAgen gel extraction kit. The mutated 9A4 scFv inserts were treated with Sfi I and Not I to prepare for ligation with the pCANTAB6 vector DNA cut with the same restriction enzymes. After ligation, the DNA mixture was ethanol precipitated and dissolved in 24 microliters of sterile deionized distilled water. Preparation of competent *E. coli* TG1 cells and electroporation was conducted as described in Example 6. Twelve separate electroporations were performed and

ultimately pooled and plated. A total of 1.5 X 10⁶ clones were obtained and stored as a glycerol stock at -80°C. A random sampling of 12 clones indicated that 9 of them (one clone failed to give sequence data) had an insert when screened by PCR, using the oligos FDTETSEQ (SEQ ID NO: 41) and pUC19R (SEQ ID NO: 39). These inserts were purified using the QIAgen kit and the sequence in the CDR3 V_H determined. The results of the sequencing data are presented below. The DNA sequence of the "Parent Sequence" is set forth in the Sequence Listing as nucleotides 383 to 409 of SEQ ID NO: 7.

10													Number of Mutations
	Parent Sequence:	5′-	GCT	AGG	GGC	GGT	AGC	CTT	GAC	TAC	TGG	-31	-
15	9A4MUT-1:	5′-	GCT	⊊ GG	GGC	GGT	AGC	CTT	GAC	TAC	⊊ GG	-3′	2
13	9A4MUT-3:	5′-	GCT	⊊ GG	<u>çc</u> c	<u>T</u> GT	A <u>T</u> C	CTT	GA <u>T</u>	TAC	TGG	-3′	6
	9A4MUT-4:	5′-	GCT	A<u>¢</u> G	GG <u>A</u>	GGT	AGC	CTT	GAC	TAC	TGG	-3′	2
20	9A4MUT-6:	5′-	G <u>T</u> T	<u>t</u> gg	GGC	GG <u>C</u>	AGC	С <u>С</u> Т	GAC	<u>C</u> AC	<u>A</u> GG	-3′	6
	9A4MUT-7:	5′-	GCT	<u>T</u> GG	GGC	GG <u>C</u>	AG <u>G</u>	<u> TA</u> T	GAC	TAC	TGG	-3'	5
	9A4MUT-8:	5′-	GCT	ANG	G <u>T</u> C	<u>a</u> gt	AGC	CTT	GAC	Т <u>С</u> С	TGG	-3'	3
25	9A4MUT-10:	5′-	GCT	A Ç G	GGC	<u>T</u> GT	AG <u>T</u>	C <u>a</u> t	GAC	TAC	<u>c</u> gc	-3′	6
	9A4MUT-12:	5′-	GCT	AGG	GG <u>T</u>	GGT	AGC	CTT	GAC	TAC	TGG	-3′	1

[0188] The mutations in the cohort above vary in number from 1 to 6 for each clone, and they occur at various positions. The mutations are indicated in bold underline in the various clones above N = nucleotide could not be assigned. Only 1 clone (9A4MUT-12) out of 8, checked in this random manner, gave parent amino acid sequence. Based on the randomness of the results shown above, a good mutated library was generated. Therefore, a biotin selection was performed to find binders to peptide 040 (SEQ ID NO: 14) which is the epitope for 9A4.

Description of Biotin selection for antibodies obtained below

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[0189] An aliquot of approximately 100 μL of mutated library stock was added to 25 mL of 2YT media containing 100 μg/mL ampicillin and 2% glucose. The culture was incubated at 37°C for approximately 60 min or until cells reached mid-log phase (OD_{600 nm} = 0.5 to 1.0). M13K07 helper phage (Pharmacia, Uppsala, Sweden) was then added to the culture to a concentration of 5 x 108 pfu/mL. The helper phage were then allowed to infect the culture for 20 min at 37°C without shaking and then for another 25 min at 37°C with shaking at 200 rpm. The infected cells were transferred to a 50 mL conical centrifuge tube and pelleted at 3000 rpm for 10 min. Cells were resuspended in 2YT media containing 100 μg/mL ampicillin and 50 μg/mL kanamycin. This culture was transferred to a fresh 250 mL flask and incubated at 30°C for 2 h during which time phage particles were produced. Cells were removed by centrifugation at 14,000 rpm for 2 min. Aliquots (1 mL) of phage were then blocked for 30 min at room temperature by the addition of PBS and NFDM to a final concentration of 1X PBS and 3% NFDM. This was accomplished by the addition of 200 μL of a 6X PBS, 18% NFDM solution to 1 mL of phage. Biotinylated peptide 040 (SEQ ID NO: 14) was added to the phage solution at concentrations ranging from 10 pM to 1 μ M. This solution was incubated for 60 min at RT. Streptavidin-coated magnetic beads (Dynal, Oslo, Norway) were blocked at RT with end-over-end shaking in 3% NFDM in PBS. Following incubation, the streptavidin beads were captured at the side of the tube with a magnet and the blocking solution carefully aspirated away. The blocked phage with the bound peptide was then added to the streptavidin beads and allowed to incubate at RT with end-over-end shaking for 15 min. Bead complexes were captured magnetically and unbound phage carefully aspirated away. Magnetic bound bead complexes were washed 4X with PBS containing 0.1% TW-20 and 4X with PBS alone. Following the final capture, bead complexes were resuspended in 100 µL of 100 mM triethylamine in PBS and neutralized with an equal volume of 1 M Tris pH 7.4. Mid-log phase E. coli TG1 cells (10 mL) were then infected with 100 μL of bead complexes. Infection was allowed to progress for 20 min at 37°C without shaking and then for another 25 min at 37°C with shaking at 200 rpm. Infected cells were then pelleted, resuspended in 500 uL of

fresh 2YT media and 500 μ L plated onto 243 x 243 mm 2YT agar plates containing 100 μ g/mL ampicillin and 2% glucose. Plates were incubated ovemight at 30°C. Following approximately 16 h of growth, the colonies were recovered by scraping and used to inoculate liquid cultures. The process was repeated for a minimum of two and a maximum of five rounds of selection.

[0190] Clones recovered from the biotin selection were grown, induced for production of scFv, which was purified according to the protocol outlined below.

Preparation of scFv mutant clones using hypotonic shock method

[0191] The culture were pelleted and resuspended in 0.8 mL ice cold TES buffer (0.2 M Tris-HCI, 0.5 nM EDTA, 0.5 M sucrose). TES (1.2 mL of ice cold 1:5 dilution) was added and the culture was incubated on ice for 30 min. The cells were pelleted at 4°C at 14,000 rpm (30 min). The scFv supernatant was added to a fresh tube containing 10 μL of 1.0 M MgCl₂. NTA-agarose (200 μL) (QlAgen) was prepared by washing in a phosphate imidazole wash buffer containing 50 mM Na phosphate, pH 8.0, 500 mM NaCl, 20 mM imidazole and 0.1% TW-20. The scFv supernatant was added to the NTA-agarose and incubated at 4°C for approximately 30 min. The NTA-agarose with scFv was spun and 500 volumes of elution buffer was added. The elution buffer consisted of 50 mM Na phosphate, pH 8.0, 500 mM NaCl and 250 mM imidazole. The eluate was vortexed and centrifuged to remove the NTA-agarose. The scFv supernatant was buffer exchanged by passing it over a NAP-5 column (Pharmacia), according to the manufacturer's instructions.

20 Measurement of off-rates for scFv constructs

[0192] Off-rates for the scFv constructs were measured by analysis of dissociation data obtained on the BIAcore (Pharmacia Biosensor) with BIAevaluation ver 2.1 software. To obtain the data on the BIAcore, streptavidin surfaces on the BIAcore chips were prepared as described previously in Example 1. Biotinylated peptide (SEQ ID NO: 14) was bound to the prepared streptavidin surfaces to RU densities ranging from about 2 - 11 RU's/surface. The lower level of derivatization would help avoid getting erroneous off-rates that could be obtained if mass transport was an issue. Purified scFv's were injected over these surfaces to allow binding of the constructs for 60 seconds. PBS buffer alone was then substituted and the dissociation of scFv was allowed to proceed for an additional 280 sec. Off-rates were calculated from these dissociation data.

Table 10 Summary of 9A4 CDR3 V_H Mutant Sequences

Amino acid sequences determined from the results of DNA sequencing. The parent sequence for clone ICAT7-1 corresponds to residues 118 to 127 of SEQ ID NO: 40.

Clone	Sequence									of	ff-rate x	10 ⁻² sec ⁻¹
ICAT7-1	С	Α	R	G	G	S	L	D	Υ	W	0.3	
15A	С	Α	R	G	Ø	R	L	D	Y	W	0.35	
16A	ပ	X	R	G	G	S	L	D	L	L	0.26	
23A	С	G	æ	G	R	S	L	D	Y	X	0.26	
24A	ပ	G	R	G	O	S	L	E	Υ	W	0.25	
26A	X	X	R	G	×	S	X	E	Y	L	0.27	
28A	X	X	R	G	O	۲	X	E	Υ	X	0.3	
9B	C	X	R	G	O	S	F	E	Y	W	0.32	
13B	C	X	R	G	G	S	X	D	F	W	0.26	
14B	C	Α	R	G	G	S	L	٥	Н	W	0.27	
20B	ပ	G	R	G	G	N	L	D	H	С	0.28	
26B	С	G	R	G	Х	T	L	E	F	W	0.34	
31B	C	G	R	G	G	ဖြ	L	D	Q	Х	0.26	
37B	C	G	R	G	G	H	L	۵	X	X	0.33	

55

30

35

40

45

38B	С	G	R	O	G	တ	ب	٥	S	C	0.26
2C	С	G	R	G	S	Ø	X	۵	Y	C	0.28
5C	С	Α	R	G	G	S	ᆫ	۵	S	W	0.25
9C	C	X	R	G	S	S	L	۵	Y	X	0.99
10C	C	G	R	G	G	S	بد	٥	Y	C	0.94
20C	C	G	R	X	G	S	X	X	뚜	C	1.29
26C	C	X	R	G	X	S	L	۵	-	X	0.95
29C	С	G	R	G	G	S	F	X	X	W	1.04
8D	C	Α	R	G	G	S	ب	۵	Z	W	1.04
16D	C	X	R	O	O	T	L	٥	Y	W	1.14
17D	C	X	X	G	R	S	L	ш	X	W	1.07
20D	С	X	R	G	X	۲	니	X	Y	W	1.02
18E	C	Α	R	G	O	S	L	D	٧	W	1.4
13G	C	G	R	G	G	S	اد	٥	7	W	1.03
17G	С	X	R	G	G	S	L	۵	щ	W	1.03
1H	C	X	R	G	G	S	L	۵	Ξ	W	1.17
4H	C	X	R	G	G	S	L	X	٧	W	1.03
1J	C	Α	R	G	G	X	_	٥	>	W	1.09
IF-5	С	Α	R	G	G	S	L	۵	Y	W	0.48

Note: Clones ICAT 7-1 and IF-5 above show CDR3 regions having the parent sequence.

[0193] It is apparent from the data presented above that changes can be made in the amino acid sequence of the parent antibody while still retaining binding to the target. While differences in the off-rate of the antibodies presented above are within an order of magnitude, by targeting different regions of the antibody V_H or V_L for mutation, or finding different antibodies obtained by immunization which have V_L and/or V_H domains derived from the same germLine V_L and V_H genes as 9A4 or 5109, one may discover antibodies with variable or enhanced binding properties relative to the parental antibodies disclosed in Examples 6 and 7 cited above.

Annex to the description

[0194]

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	-404	0 > 32	,													
					. 1 .		~ 1~	0	T1-	a1-	.1-	~1 -	T1 -	~ 1		
15		Leu	mec	MIG		Ala	GIII	ser	116		AIA	GIN	TTE	GIN	Leu	vaı
	1				5					10					15	
	- 1-	•	~ 1		63 .	• -		_	_	-1				_		_
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									-							
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	35 40 45
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	50 55 60
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primer for 5' 9A4 VL region, including an Nco I restriction endonuclease site.

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                                           25
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51

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	_		_				_	_	_	_		_				
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	65					70					75					80
25	Leu (G] n	Wat	han	7~~	Lau	Tua	Co	C 111	7.00	Co=	<i>a</i> 1	Wat	M	(Th. 222	۵.,
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					0,5					90					33	
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	JJ J.	J. J	•
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25
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                                                                              18
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25	Ala	Gln	Pro	Ala	Met	Ala	Glu	Val	Gln	Leu	Val	Glu	Ser	Gly	Gly	Gly
				20					25					30		_
	Ser	Val	Gln	Pro	Gly	Gly	Ser	Leu	Lys	Leu	Ser	Сув	Ala	Ala	Ser	Gly
30			35					40					45			
				_		_			_	_					_	_
	Phe		Phe	Asn	Thr	Tyr		Met	Ser	Trp	Val		Gln	Thr	Pro	Asp
		50					55					60				
35	Lvs	Arg	Leu	Glu	Tro	Val	Ala	Thr	Ile	Asn	Ser	Asn	Glv	Glv	Leu	Thr
	65					70					75			2		80
	Phe	Tyr	Ala	Asp	Ser	Val	Lys	Gly	Arg	Phe	Thr	Ile	Ser	Arg	Asp	Asn
40					85					90					95	
40																
	Ala	Lys	Asn	Thr	Leu	Tyr	Leu	Gln	Met	Asn	Arg	Leu	Lys	Ser	Gly	Asp
				100					105					110		
45	_			_	_					_			_		_	_
•5	Ser	GIÀ		Tyr	Tyr	Cys	Val		GIY	Tyr	Ser	Asn		Ala	Arg	Trp
			115					120					125			
	G]v	Gln	Glv	Ala	Lev	Val	Thr	Val	Ser	Ser	G) v	G) v	G) v	G) v	Ser	Glv
50	,	130	1				135				1	140	1	1		1
7U																
	Gly	Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Ser	Asp	Val	Val	Met	Thr	Gln
	145					150					155					160
5 <i>5</i>								_								
	Th-	Dva	T 411	Thr	T 011	00-	37-3	Th-	T1-	01.4	~ I ~	C	21-	C ~ ~	T 3 -	Ca-

		165	170		175
5	Cys Lys Ser Ser 180	Gln Ser Le	u Leu Gly Ser 185		Thr Tyr Leu 190
	Ile Trp Leu Leu 195	Gln Arg Pr	o Gly Gln Ser 200	Pro Lys Arg I 205	Seu Ile Phe
10	Leu Val Ser Glu 210	Leu Asp Se	=	Asp Arg Phe 1 220	Thr Gly Ser
15	Gly Ser Gly Thr 225	Asp Phe Th	r Leu Lys Ile	Ser Arg Ala 0 235	Glu Ala Glu 240
20	Asp Leu Gly Val	Tyr Tyr Cy 245	s Cys Gln Gly 250	Thr His Phe H	Pro His Thr 255
	Phe Gly Ala Gly 260	Thr Lys Le	u Glu Leu Lys 265		Glu Gln Lys 270
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                                                                             62
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                                                10
                                                                     15
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              1
                               5
                                                   10
                                                                         15
30
            Glu Xaa Gly Val Ser Tyr
                          20
```

Claims

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A method for monitoring biological media for protein fragments which comprises;

contacting said biological media with a capture antibody; said capture antibody being active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 1 and 2; and

contacting said biological media with a detection antibody; said capture antibody being active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 3 and 4; and

detecting the amount of collagen fragments bound to said capture and detection antibodies; or

contacting said biological media with a capture antibody; said capture antibody being active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 3 and 4; and

contacting said biological media with a detection antibody; said detection antibody being active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 1 and 2; and

- detecting the amount of collagen fragments bound to said capture and detection antibodies.
- A method of claim 1 wherein said protein fragments are collagen fragments generated by collagenase cleavage of articular cartilage.

- 3. A method of claim 2 wherein said protein fragments are generated from collagenase cleavage of type II collagen.
- 4. A method for monitoring biological media for protein fragments which comprises;
- 5 contacting said biological media with an antibody active against the sequences set forth in the Sequence Listing as SEQ ID NOS: 3 and 4; and
 - detecting the amount of protein fragments bound to said antibody.
- 10 5. A method of claim 4 wherein said protein fragments are collagen fragments.

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- 6. A method of claim 1 wherein said capture and detection antibodies are monoclonal antibodies.
- 7. A method of claim 2 wherein said capture and detection antibodies are monoclonal antibodies.

8. A method of claim 1 wherein said capture antibody is a genetically engineered antibody.

- 9. A method of claim 1 wherein said detection antibody is a genetically engineered antibody.
- 20 10. A method of claim 1 wherein said capture antibody, designated 9A4, has the V_H sequence as set forth in the Sequence Listing as SEQ ID NOS: 5 and 32 and the V_L sequence as set forth in the Sequence Listing as SEQ ID NOS: 6 and 33
- 11. A method of claim 1 wherein said detection antibody, designated 5109, has the V_H sequence as set forth in the Sequence Listing as SEQ ID NOS: 10 and 48 and the V_L sequence as set forth in the Sequence Listing as SEQ ID NOS: 11 and 49.
 - 12. A method of claim 4 wherein said antibody is a monoclonal antibody.
- 30 13. A method of claim 4 wherein said antibody is a genetically engineered antibody.
 - 14. A method of claim 4 wherein said antibody, designated 5109, has the V_H sequence as set forth in the Sequence Listing as SEQ ID NOS: 10 and 48 and the V_L sequence as set forth in the Sequence Listing as SEQ ID NOS: 11 and 49.
 - 15. The antibody, designated 9A4, having the V_H sequence as set forth in the Sequence Listing as SEQ ID NOS: 5 and 32 and the V_L sequence as set forth in the Sequence Listing as SEQ ID NOS: 6 and 33.
- 16. The antibody, designated 5109, having the V_H sequence as set forth in the Sequence Listing as SEQ ID NOS: 10 and 48 and the V_L sequence as set forth in the Sequence Listing as SEQ ID NOS: 11 and 49.
 - 17. A cell line that produces a specific binding partner that binds to peptides consisting essentially of the structures as set forth in the Sequence Listing as SEQ ID NOS: 1 or 2, the cell line having the identifying characteristics of ATCC HB-12436.
 - 18. A cell line that produces a specific binding partner that binds to peptides consisting essentially of the structures as set forth in the Sequence Listing as SEQ ID NOS: 3 or 4, the cell line having the identifying characteristics of ATCC HB 12435.
- 50 19. A genetically engineered form of antibody 9A4 such as that expressed by E. coli p9A41CAT7-1 or p9A41F-5 having the characteristics of ATCC-98593 and ATCC-98592, respectively.
 - 20. A genetically engineered form of antibody 5109 such as that derived from E. coli p5109CscFv7, which may be substituted for 5109, and having the characteristics of ATCC-98594.
 - 21. A bispecific antibody produced by hybridization of the antibodies 9A4 and 5109 wherein each half antibody recognizes its respective binding partner.

- 22. A bispecific antibody which is a genetically engineered combination of antibodies 9A4 and 5109 produced by combining the V_L and V_H domains of the two antibodies in the form V_L (5109)-linker-V_H (5109)-linker-V_L (9A4)-linker-V_L V_H(9A4) and equivalents thereof.
- 5 23. A method for monitoring collagen fragments in biological media which comprises: contacting said biological media with a bispecific antibody 9A4/5109 of claim 22; and detecting the amount of collagen fragments bound to said antibody.
 - 24. A bispecific antibody of claim 22 produced from genetically modified variants of antibodies 5109 and 9A4.
 - 25. An antibody related to the antibodies 9A4 or 5109 by virtue of the V_L and V_H genes being derived from the same germline genes as the corresponding V_L and V_H genes of 9A4 and 5109 and where changes occur in the sequence, but where the V_L - V_H variant combinations maintain the binding properties of antibody 9A4 or 5109.
- 15 26. An antibody of claim 1 which is labeled to facilitate detection.

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- 27. An antibody of claim 26 wherein said label is radioactive, optical, enzymatic, fluorescent polarizing or fluorescent quenching.
- 20 28. An antibody of claim 1 which is labeled to facilitate capture of said antibody.
 - 29. An antibody of claim 28 wherein said label is biotin or magnetic particles.

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